

### **PCT**

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau

### INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:			(11) International Publication Number: WO 99/43794				
C12N 9/28, A21D 8/04		A1	(43	3) International	Publication	Date:	2 September 1999 (02.09.99)
(21) International Application Number: PC	T/DK99/000		38	(81) Designated BY, CA	States: AL	C, AU, AZ, BA, BB, BG, BR, DE, DK, EE, ES, FI, GB, GD,	

26 February 1999 (26.02.99)

(30) Priority Data: 27 February 1998 (27.02.98) DK 60/077,795 12 March 1998 (12.03.98)

(71) Applicant (for all designated States except US): NOVO NORDISK A/S [DK/DK]; Novo Allé, DK-2880 Bagsværd (DK).

(72) Inventors; and

(22) International Filing Date:

(75) Inventors/Applicants (for US only): CHERRY, Joel, Robert [US/US]; 916 Anderson Road, Davis, CA 95616 (US). SVENDSEN, Allan [DK/DK]; Bakkeledet 28, DK-3460 Birkerød (DK). ANDERSEN, Carsten [DK/DK]; Højeloft Vænge 162, DK-3500 Værløse (DK). BEIER, Lars [DK/DK]; Skeltoftevej 16, st. h, DK-2800 Lyngby (DK). FRANDSEN, Torben, Peter [DK/DK]; Alhambravej 22, 1, th, DK-1826 Frederiksberg C (DK).

(74) Common Representative: NOVO NORDISK A/S; Corporate Patents, Novo Allé, DK-2880 Bagsværd (DK).

GE, GH, GM, HR, HU, ID, IL, IN, IS, IP, KE, KG, KP, GE, GH, GM, HR, HU, ID, IL, IN, IS, IP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG) SN, TD, TG).

#### Published

With international search report.

(54) Title: MALTOGENIC ALPHA-AMYLASE VARIANTS

#### (57) Abstract

The inventors have modified the amino acid sequence of a maltogenic alpha-amylase to obtain variants with improved properties, based on the three-dimensional structure of the maltogenic alpha-amylase Novamyl. The variants have altered physicochemical properties., e.g. an altered pH optimum, improved thermostability, increased specific activity, an altered cleavage pattern or an increased ability to reduce retrogradation of starch or staling of bread.

### FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Pinland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LÜ	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BI	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL.	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

1/PR15

10/069908 JC19 Racid PCT/PTO 28 FEB 2002

WO 99/43794

30

PCT/DK99/00088

1

### **MALTOGENIC ALPHA-AMYLASE VARIANTS**

### **FIELD OF THE INVENTION**

The present invention relates to variants of maltogenic amylase and to methods of constructing such variants.

### 5 BACKGROUND OF THE INVENTION

Maltogenic alpha-amylase (glucan 1,4- $\alpha$ -maltohydrolase, E.C. 3.2.1.133) is able to hydrolyze amylose and amylopectin to maltose in the alpha-configuration, and is also able to hydrolyze maltotriose as well as cyclodextrin.

A maltogenic alpha-amylase from *Bacillus* (EP 120 693) is commercially available under the trade name Novamyl® (product of Novo Nordisk A/S, Denmark) and is widely used in the baking industry as an anti-staling agent due to its ability to reduce retrogradation of starch. Novamyl® shares several characteristics with cyclodextrin glucanotransferases (CGTases), including sequence homology (Henrissat B., Bairoch A. 1996) and formation of transglycosylation products (Christophersen, C., et al., 1997, Starch, vol. 50, No. 1, 39-45).

Cyclomaltodextrin glucanotransferase (E.C. 2.4.1.19), also designated cyclodextrin glucanotransferase or cyclodextrin glycosyltransferase, abbreviated herein as CGTase, catalyses the conversion of starch and similar substrates into cyclomaltodextrins via an intramolecular transglycosylation reaction, thereby forming cyclomaltodextrins (or CD) of various sizes.

CGTases are widely distributed and from several different bacterial sources, Clostridium, Corynebacterium, Bacillus. Brevibacterium, including Thermoanaerobacter and Thermoanaerobacterium have Micrococcus. extensively described in the literature. A CGTase produced by Thermoanaerobacter sp. 25 has been reported in Norman B E, Jørgensen S T; Denpun Kagaku 1992 39 99-106, and WO 89/03421, and the amino acid sequence has been disclosed in WO 96/33267. The sequence of CGTases from Thermoanaerobacterium thermosulfurigenes and from Bacillus circulansis available on the Internet (SCOP or PDF home pages) as pdf file 1CIU, and the sequence of a CGTase from B. circulans is available as pdf file 1CDG.

Tachibana, Y., Journal of Fermentation and Bioengineering, 83 (6), 540-548 (1997) describes the cloning and expression of a CGTase. Variants of CGTases have been described by Kim, Y. H., Biochemistry and Molecular Biology International, 41 (2), 227-234 (1997); Sin K-A, Journal of Biotechnology, 32 (3), 283-288 (1994); D Penninga, Biochemistry, 34 (10), 3368-3376 (1995); and WO 96/33267.

Recently, the tertiary structure of several CGTases have been reported. Hofman et al. [Hofman B E, Bender H, Schultz G E; <u>J. Mol. Biol.</u> 1989 **209** 793-800] and Klein & Schulz [Klein C, Schulz G E; J. Mol. Biol. 1991 217 737-750] report the tertiary structure of a CGTase derived from Bacillus circulans Strain 8, Kubota et al. [Kubota M, 5 Matsuura Y, Sakai S and Katsube Y; Denpun Kagaku 1991 38 141-146] report the tertiary structure of a CGTase derived from Bacillus stearothermophilus TC-91, Lawson et al. [Lawson C L, van Montfort R, Strokopytov B, Rozeboom H J, Kalk K H, de Vries G E. Penninga D, Dijkhuizen L, and Dijkstra B W; J. Mol. Biol. 1994 236 590-600] report the tertiary structure of a CGTase derived from Bacillus circulans Strain 251, 10 Strokopytov et al. [Strokopytov B, Penninga D, Rozeboom H J; Kalk K H, Dijkhuizen L and Dijkstra B W; Biochemistry 1995 34 2234-2240] report the tertiary structure of a CGTase derived from Bacillus circulans Strain 251, which CGTase has been complexed with acarbose, an effective CGTase inhibitor, and Knegtel et al. [Knegtel R M A, Wind R D, Rozeboom H J, Kalk K H, Buitelaar R M, Dijkhuizen L and Dijkstra B 15 W; J. Mol. Biol. 1996 256 611-622] report the tertiary structure of a CGTase derived from Thermoanaerobacterium thermosulfurigenes.

### **BRIEF DISCLOSURE OF THE INVENTION**

The inventors have modified the amino acid sequence of a maltogenic alphaamylase to obtain variants with improved properties, based on the three-dimensional structure of the maltogenic alpha-amylase Novamyl. The variants have altered physicochemical properties., e.g. an altered pH optimum, improved thermostability, increased specific activity, an altered cleavage pattern or an increased ability to reduce retrogradation of starch or staling of bread.

Accordingly, the present invention provides a method of constructing a variant of a parent maltogenic alpha-amylase, wherein the variant has at least one altered property as compared to said parent maltogenic alpha-amylase, which method comprises:

- i) analyzing the structure of the maltogenic alpha-amylase to identify, on the basis of an evaluation of structural considerations, at least one amino acid residue or at least one structural region of the maltogenic alpha-amylase, which is of relevance for altering said property;
  - ii) constructing a variant of the maltogenic alpha-amylase, which as compared to the parent, has been modified in the amino acid residue or structural part identified in i) so as to alter said property; and
    - iii) testing the resulting maltogenic alpha-amylase variant for said property.

The property which may be altered by the above methods of the present invention may be, e.g., stability, pH dependent activity, ability to reduce retrogradation

of starch or staling of bread, specific activity, or substrate specificity. Thus, the variant may have, e.g., increased thermostability or higher activity at a lower pH an altered pH optimum, improved thermostability, increased specific activity or increased ability to reduce retrogradation of starch or staling of bread

In still further aspects the invention relates to variants of a maltogenic alphaamylase, the DNA encoding such variants and methods of preparing the variants. Finally, the invention relates to the use of the variants for various industrial purposes, in particular baking.

### **DETAILED DISCLOSURE OF THE INVENTION**

### 10 Maltogenic alpha-amylase

25

The maltogenic alpha-amylase is an enzyme classified in EC 3.2.1.133. The enzymatic activity does not require a non-reducing end on the substrate and the primary enzymatic activity results in the degradation of amylopectin and amylose to maltose and longer maltodextrins. It is able to hydrolyze amylose and amylopectin to maltose in the alpha-configuration, and is also able to hydrolyze maltotriose as well as cyclodextrin.

A particularly preferred maltogenic alpha-amylase is the amylase cloned from Bacillus as described in EP 120 693 (hereinafter referred to as Novamyl). Novamyl has the amino acid sequence set forth in amino acids 1-686 of SEQ ID NO: 1. Novamyl is encoded in the gene harbored in the Bacillus strain NCIB 11837 which has the nucleic acid sequence set forth in SEQ ID NO:1. The three-dimensional structure of Novamyl is described below.

In general, a preferred maltogenic alpha-amylase should have one or more of the following properties:

- i) a three dimensional structural homology to Novamyl,
- ii) an amino acid sequence having at least 70 % identity to SEQ ID NO: 1, preferably at least 80 % or 90 %, e.g. 95 % or 98 %,
- iii) a DNA sequence which hybridizes to the DNA sequence set forth in SEQ ID NO:1 or to the DNA sequence encoding Novamyl harbored in the *Bacillus* strain NCIB 11837;
- iv) a calcium binding site comprising a coordination equivalent to a backbone carbonyl atom from Asn77, sidechain atom OE2 and OE1 from Glu102, a sidechain atom OD1 from Asp79, a sidechain atom OD1 from Asp76, and a sidechain atom OE1 from Glu101, and one water molecule WAT V21, atom OW0, wherein the positions are as shown in Appendix 1;

v) a sequence of five amino acids corresponding to Pro-Ala-Gly-Phe-Ser in a position equivalent to residues 191-195 in the amino acid sequence shown in SEQ ID NO: 1; and

The structural homology referred to above in i) is based on other sequence bornologies, hydrophobic cluster analysis or by reverse threading (Huber, T; Torda, AE, PROTEIN SCIENCE Vol. 7, No. 1 pp. 142-149 (1998)) and which by any of these methods is predicted to have the same tertiary structure as Novamyl, wherein the tertiary structure refers to the overall folding or the folding of Domains A, B, and C, more preferably including Domain D, and most preferably including Domain E. Alternatively, a structural alignment between Novamyl and a maltogenic alpha-amylase may be used to identify equivalent positions.

The calcium binding site referred to above in iv) is based on a calcium binding site identified in the three-dimensional structure of Novamyl, and is discussed below in the section "Calcium binding sites."

The "equivalent position" referred to above in v) is based on amino acid or DNA sequence alignment or structural homology using methods known in the art.

### Three-dimensional structure of maltogenic alpha-amylase

Novamyl was used to elucidate the three-dimensional structure forming the basis for the present invention.

The structure of Novamyl was solved in accordance with the principle for x-ray crystallographic methods, for example, as given in X-Ray Structure Determination, Stout, G.K. and Jensen, L.H., John Wiley & Sons, Inc. NY, 1989.

The structural coordinates for the solved crystal structure of Novamyl at 2.2 Å resolution using the isomorphous replacement method are given in standard PDB format (Protein Data Bank, Brookhaven National Laboratory, Brookhaven, CT) as set forth in Appendix 1. It is to be understood that Appendix 1 forms part of the present application. In the context of Appendix 1, the following abbreviations are used: CA refers to calcium ion or alpha-carbon atom of the polypeptide backbone, WAT refers to water or to calcium, MAL refers to maltose, HEX refers to a carbohydrate unit of a substrate analogue, and SUL refers to a sulfate ion.

Amino acid residues of the enzyme are identified herein by their respective one- or three-letter amino acid code.

The structure of said maltogenic alpha-amylase is made up of five globular domains, ordered A, B, C, D and E. The domains can be defined as being residues 1-35 132 and 204-403 for Domain A, residues 133-203 for Domain B, residues 404-496 for Domain C, residues 497-579 for Domain D, and residues 580-686 for Domain E,

wherein the numbering refers to the amino acid sequence in SEQ ID NO: 1. Features of Domains A, B, and C of particular interest are described below.

### Domain A

Domain A is the largest domain and contains the active site which comprises a cluster of three amino acid residues, D329, D228 and E256, spatially arranged at the bottom of a cleft in the surface of the enzyme. The structure of Domain A shows an overall fold in common with the α-amylases for which the structure is known, viz. the (beta/alpha) 8 barrel with eight central beta strands (numbered 1-8) and eight flanking a-helices. The β-barrel is defined by McGregor op. cit. The C-terminal end of the beta strand 1 is connected to helix 1 by a loop denoted loop 1 and an identical pattern is found for the other loops, although the loops show some variation in size and some can be quite extensive.

The eight central beta-strands in the (beta/alpha) 8 barrel superimpose reasonably well with the known structures of CGTases. This part of the structure, including the close surroundings of the active site located at the C-terminal end of the beta-strands, shows a high degree of identity with CGTases.

In contrast, the loops connecting the beta-strands and alpha helices display a high degree of variation from the known structures of CGTases. These loops constitute the structural context of the active site, and the majority of the contacts to the substrate is found among residues located in these loops. Distinguishing characteristics such as substrate specificity, substrate binding, pH activity profile, substrate cleavage pattern, and the like, are determined by specific amino acids and the positions they occupy in these loops. In Novamyl Domain A contains two calcium binding sites, one of which is homologous to the calcium binding site in CGTases; the other is unique to Novamyl.

The structure of the calcium binding site is discussed further below in the section "Calcium binding sites."

### Domain B

Domain B, also referred to as loop 3 of the (beta/alpha) 8 barrel, in comprises amino acid residues 133-203 of the amino acid sequence shown in SEQ ID NO: 1. The structure is partially homologous to the structure of Domain B in CGTases, the most striking difference being the presence of a five amino acid insert corresponding to positions 191-195 in the amino acid sequence shown in SEQ ID NO: 1 which is not found in the CGTases. This insert is spatially positioned close to the active site residues and in close contact to the substrate.

### Domain C

15

20

Domain C in Novamyl comprises amino acid residues 404-496 of the amino acid sequence shown in SEQ ID NO: 1. Domain C is composed entirely of β-strands which form a single 8-stranded sheet structure that folds back on itself, and thus may be described as a β-sandwich structure. One part of the β-sheet forms the interface to Domain A.

### Calcium binding sites

The structure of the maltogenic alpha-amylase exhibits three calcium-binding sites; that is, three calcium ions are found to be present in the structure. In common with most of the known family 13 structures, one calcium ion, WAT 693 in Appendix 1, is located between the A and B domains. This calcium ion is coordinated by a backbone carbonyl atom from Gln184 and His232, sidechain atoms OD2 and OD1 from Asp198, a sidechain atom OD1 from Asn131, and three water molecules WAT V1, WAT V5 and WAT V8.

A second calcium ion is located in the A domain and is common to CGTases, but not found in  $\alpha$ -amylases. The calcium ion WAT 694 is coordinated by a backbone carbonyl atom from Gly48 and Asp23, sidechain atom OD2 from Asp50, a sidechain atom OD1 from Asp21, a sidechain atom OD1 from Asp26, and a sidechain atom OD1 from Asp27, and one water molecule WAT V62.

The third calcium ion is located in the A Domain and is unique to Novamyl. The calcium ion is WAT 692 and the coordination comprises a backbone carbonyl atom from Asn77, sidechain atom OE2 and OE1 from Glu102, a sidechain atom OD1 from Asp79, a sidechain atom OD1 from Asp76, and a sidechain atom OE1 from Glu101, and one water molecule WAT V21.

### 25 Substrate Binding Site

Parts of the loop discussed above in the context of domains A and B are of particular interest for substrate interaction and active site reactivity. In particular, in domain A, residues 37-45 in loop 1, residues 261-266 in loop 5, residues 327-330 in loop 7 and residues 370-376 in loop 8; in domain B, residues 135-145 in loop 3, residues 173-180 and 188-196 in loop 3, wherein residue positions correspond to the amino acids in the amino acid sequence in SEQ ID NO: 1.

Without being limited to any theory, it is presently believed that binding between a substrate and an enzyme is supported by favorable interactions found within a sphere of 4 to 6 Å between the substrate molecule and the enzyme, such as hydrogen bonds and/or strong electrostatic interaction. The following residues of

25

35

Novamyl (SEQ ID NO: 1), are within a distance of 6 Å of the substrate HEX and thus believed to be involved in interactions with said substrate:

44, 89, 90, 92, 93, 127, 129, 132, 135, 177, 178, 188, 191, 194, 196, 226, 228, 229, 230, 231, 232, 256, 258-261, 288, 328, 329, 371, 372, 373, 376, and 690.

The following residues of Novamyl are within a distance of 4 Å of the substrate HEX and thus believed to be involved in interactions with said substrate:

90, 92, 93, 129, 132, 177, 188, 189, 190, 191, 196, 226, 228, 229, 231, 232, 256, 258, 259, 260, 261, 328, 329, 372, 376, and 690.

# Homology building of Novamyl®

The structure of the Novamyl® was model built on the structure disclosed in Appendix 1 herein. The structure of other maltogenic alpha-amylases may be built analogously.

A model structure of a maltogenic alpha-amylase can be built using the Homology program or a comparable program, eg., Modeller (both from Molecular Simulations, Inc., San Diego, CA). The principle is to align the sequence of the maltogenic alpha-amylase with the known structure with that of the maltogenic alpha-amylase for which a model structure is to be constructed. The structurally conserved regions can then be built on the basis of consensus sequences. In areas lacking homology, loop structures can be inserted, or sequences can be deleted with subsequent bonding of the necessary residues using, e.g., the program Homology. Subsequent relaxing and optimization of the structure should be done using either Homology or another molecular simulation program, e.g., CHARMm from Molecular Simulations.

# Methods for designing novel maltogenic alpha-amylase variants

In a first aspect, the invention relates to a method of constructing a variant of a parent maltogenic alpha-amylase, wherein said variant has at least one altered property as compared to said parent  $\alpha$ -amylase, which method comprises:

- i) analyzing the structure of the maltogenic alpha-amylase to identify at least one amino acid or structural region of said α-amylase, which, on the basis of structural or functional considerations, is determined to be of relevance for altering said property of the parent maltogenic alpha-amylase;
  - ii) constructing a variant of the maltogenic alpha-amylase, which as compared to the parent, has been modified in the amino acid residue or structural region identified in i) has been modified so as to alter said property; and
    - iii) testing the resulting variant for said property.

35

The structural part which is identified in step i) of the method of the invention may be composed of one amino acid residue. However, normally the structural part comprises more than one amino acid residue, typically constituting one of the above parts of the maltogenic alpha-amylase structure such as one of the A, B, C, D or E domains, an interface between any of these domains, a calcium binding site, a loop structure, the substrate binding site, or the like.

The structural or functional considerations may involve an analysis of the relevant structure or structural part and its contemplated impact on the function of the enzyme. For example, an analysis of the functional differences between maltogenic alpha-amylase and the various CGTases may be used for assigning certain properties of Novamyl to certain parts of the Novamyl structure or to contemplate such relationship. For instance, differences in the pattern or structure of loops surrounding the active site may result in differences in access to the active site of the substrate and thus differences in substrate specificity and/or cleavage pattern.

Furthermore, parts of a maltogenic alpha-amylase involved in substrate binding, and thus, for example, substrate specificity and/or cleavage, calcium ion binding, important, for example, for the calcium dependency of the enzyme, and the like, have been identified (*vide infra*).

The modification of an amino acid residue or structural region is typically accomplished by suitable modifications of a DNA sequence encoding the parent enzyme in question. The modification may be substitution, deletion or insertion of an amino acid residue or a structural part.

The property to be modified may be stability (e.g. thermostability), pH dependent activity, substrate specificity, specific activity or ability to reduce retrogradation of starch or staling of bread. Thus, the altered property may be an altered specific activity at a given pH and/or an altered substrate specificity, such as an altered pattern of substrate cleavage or an altered pattern of substrate inhibition.

In step ii) of the method according to the invention the part of the structure to be identified is preferably one which in the folded enzyme is believed to be in contact with the substrate (cf, the disclosure above in the section entitled "Substrate Binding Site") or involved in substrate specificity and/or cleavage pattern, and/or one which is in contact with one of the calcium ions and/or one, which is contributing to the pH or temperature profile of the enzyme, or is otherwise responsible for the properties of the maltogenic alpha-amylase.

Described in the following are specific types of variants which have been designed by use of the method of the invention.

The variants of the invention may comprise additional modifications in addition to the modifications described herein. The variants preferably have an amino acid



having more than 70 % identity with SEQ ID NO: 1, preferably more than 80 %, particularly more than 90 %, especially more than 95 %, e.g. more than 98 %.

### Maltogenic alpha-amylase variants with altered pH dependent activity profile

The pH dependent activity profile can be changed by changing the pKa of residues within 10 Å of the active site residues of the maltogenic alpha-amylase. Changing the pKa of the active site residues is achieved, e.g., by changing the electrostatic interaction or hydrophobic interaction between functional groups of amino acid side chains of a given amino acid residue and its close surroundings. To obtain a higher activity at a higher pH, negatively charged residues are placed near a hydrogen donor acid, whereas positively charged residues placed near a nucleophilic acid will result in higher activity at low pH. Also, a decrease in the pKa can be obtained by reducing the accessibility of water or increasing hydrophobicity of the environment.

Thus, another aspect of the present invention relates to a variant of a parent maltogenic alpha-amylase, in which the variant has an altered pH dependent activity profile as compared to the parent, wherein the variant may be obtained by the following method:

- i) identifying an amino acid residue within 15 Å from an active site residue of a maltogenic alpha-amylase in the three-dimensional structure of said parent maltogenic alpha-amylase, in particular 10 Å from an active site residue, wherein said amino acid residue is contemplated to be involved in electrostatic or hydrophobic interactions with an active site residue;
- ii) substituting, in the structure, said amino acid residue with an amino acid residue which changes the electrostatic and/or hydrophobic surroundings of an active site residue, and evaluating the accommodation of the amino acid residue in the 25 structure,
  - iii) optionally repeating step i) and/or ii) recursively until an amino acid substitution has been identified which is accommodated into the structure,
  - iv) constructing a maltogenic alpha-amylase variant resulting from steps i) and ii), and optionally iii), and testing the pH dependent enzymatic activity of said variant.

In a preferred embodiment, the variant of a maltogenic alpha-amylase having an altered pH dependent activity profile as compared to the parent maltogenic alpha-amylase comprises a modification of an amino acid residue corresponding to one or more of the following residues of the amino acid sequence set forth in SEQ ID NO: 1:

D127, V129, F188, A229, Y258, V281, F284, T288, N327, M330, G370, N371, and D372,

L71, S72, V74, L75, L78, T80, L81, G83, T84, D85, N86, T87, G88, Y89, H90, G91, T94, R95, D96, F97, I174, S175, N176, D178, D179, R180, Y181, E182, A183, Q184, K186, N187, F188, T189, D190, A192, G193, F194, S195, L196.

In more preferred embodiment, the variant comprises a modification 5 corresponding to one or more of the following modifications in the amino acid sequence set forth in SEQ ID NO: 1:

A229S/T/G/V, D127N/L, V129S/T/G/V, F188E/K/H. Y258E/D/K/R/F/N, T288E/K/R, N327D. M330L/F/I/D/E/K, F284K/H/D/E/Y, G370N. V281L/T. N371D/E/G/K, and D372N/V,

L711, S72C, V741, L75N/D/Q/I/V, L78N/I, T80I/L/V/S/N/G, L81I/V/S/T/N/Q/K/H, G83A/S/T/N/Q/E/D/R/H/L, T84S/A/N/D/G, D85A/T/S/N/G, N86Q/E/D/Y/H/K, T87S/I, G88A/S/T, Y89F, H90N/Q/K, G91A/S/T, T94N/D/A/M/V/I, R95K/Q, D96N/V/Q/I, F97Y, 1174N/Q/L, S175T/A/N/D, N176S/T/H/Q/P, D178N/Q/E/K/H, D179Y/N/H, R180W, Y181R/F/C/L. E182D. A183S/C/G, Q184E, K186R. N187Q/E/L/F/H/K/V/L, 15 F188Y/L/I/H/N, T189N/D/A/S/H/Y/G, D190E/Q/H/N/K, A192T/D/E/N/K, G193A/S/T, F194Y, S195N/D/E/R/K/G, L1961.

Similar modifications may be introduced in equivalent positions of other maltogenic alpha-amylases. Variants of particular interest have a combination of one or more of the above with any of the other modifications disclosed herein.

### 20 Maltogenic alpha-amylase variants with altered stability

A variant with improved stability (typically increased stability) may be obtained by stabilization of calcium binding, substitution with proline, substitution of histidine with another amino acid, introduction of an interdomain disulfide bond, removal of a deamidation site, altering a hydrogen bond contact, filling in an internal structural cavity 25 with one or more amino acids with bulkier side groups, introduction of interdomain interactions, altering charge distribution, helix capping, or introduction of a salt bridge.

### Calcium binding

The invention provides a variant of a parent maltogenic alpha-amylase, which has an altered stability due to an altered stabilization of calcium (Ca2+) binding. The 30 enzyme variant may have altered thermostability or pH dependent stability, or it may have maltogenic alpha-amylase activity in the presence of a lower concentration of calcium ion. It is presently believed that amino acid residues located within 10 Å from a calcium ion are involved in or are of importance for the Ca2+ binding capability of the enzyme.

25

The amino acid residues found within a distance of 10 Å from the Ca<sup>2+</sup> binding sites of the maltogenic alpha-amylase with the amino acid sequence set forth in SEQ ID NO: 1 were determined as described in Example 2 and are as follows:

16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28 29, 30, 31, 32, 33,35, 36, 40, 5 46, 47, 48, 49, 50, 51, 52, 53, 54, 56, 73, 74, 75, 76, 77, 78, 79, 80, 81, 87, 88, 89, 91, 93, 94, 95, 96, 99, 100, 101, 102, 103, 104, 105, 109, 129, 130, 131, 132, 133, 134, 145, 150, 167, 168, 169, 170, 171, 172, 174, 177, 180, 181, 182, 183, 184, 185, 186, 187, 188, 189, 196, 197, 198, 199, 200, 201, 202, 206, 210, 228, 229, 230, 231, 232, 233, 234, 235, 237, 378, and 637.

In order to construct a variant according to this aspect of the invention it is desirable to substitute at least one of the above mentioned amino acid residues, which is determined to be involved in a non-optimal calcium binding, with any other amino acid residue which improves the Ca2+ binding affinity of the variant enzyme. Accordingly, another aspect of the invention relates to a method of constructing a 15 variant of a parent maltogenic alpha-amylase wherein said variant has a stabilised Ca2+ binding as compared to said parent, which method comprises:

- i) identifying an amino acid residue within 10 Å from a Ca2+ binding site of a maltogenic alpha-amylase in a model of the three-dimensional structure of said  $\alpha$ amylase which, from structural or functional considerations, is determined to be 20 responsible for a non-optimal calcium ion interaction;
  - ii) constructing a variant in which said amino acid residue is substituted with another amino acid residue which, from structural or functional considerations, is determined to be important for establishing an altered Ca2+ binding affinity; and
    - iii) testing the Ca2+ binding of the resulting maltogenic alpha-amylase variant.

Substituting an amino acid residue responsible for non-optimal calcium ion interaction with another residue may alter a calcium ion binding interaction of the enzyme. For instance, the amino acid residue in question may be selected on the basis of one or more of the following objectives:

- a) to obtain an improved interaction between a calcium ion and an amino acid 30 residue as identified from the structure of the maltogenic alpha-amylase. For instance, if the amino acid residue in question is exposed to a surrounding solvent, it may be advantageous to increase the shielding of said amino acid residue from the solvent so as to stabilize the interaction between said amino acid residue and a calcium ion. This can be achieved by substituting said residue, or an amino acid residue in the vicinity of 35 said residue contributing to the shielding, with an amino acid residue with a bulkier side group or which otherwise results in an improved shielding effect.
  - b) to stabilize a calcium binding site, for instance by stabilizing the structure of the maltogenic alpha-amylase, e.g. by stabilizing the contacts between two or more of

15

20

the five domains or stabilizing one or more of the individual domains as such. This may, e.g., be achieved by providing for a better coordination to amino acid side chains, which may, e.g., be obtained by substituting an N residue with a D residue and/or a Q residue with an E residue, e.g. within 10 Å, and preferably within 3 or 4 Å, of a calcium binding site.

c) to improve the coordination between the calcium ion and the calcium binding residues, e.g., by improving the interaction between the ion and the coordinating residues or increasing the number of sidechain coordinations by substituting a coordinating water with an amino acid sidechain.

d) replace water by a coordinating calcium amino acid residue.

Preferably, the amino acid residue to be modified is located within 8 Å of a Ca<sup>2+</sup> ion, preferably within 5 Å of a Ca<sup>2+</sup> ion. The amino acid residues within 8 Å and 5 Å, respectively, may easily be identified by an analogous method used for identifying amino acid residues within 10 Å (cf. Example 2).

In a preferred embodiment, the variant of a maltogenic alpha-amylase having an altered Ca<sup>2+</sup> binding as compared to the parent maltogenic alpha-amylase comprises a substitution of an amino acid residue corresponding to one or more of the following residues of the amino acid sequence set forth in SEQ ID NO: 1:

D17, A30, S32, R95, H103, N131, Q201, I174, and/or H169,

V74, L75, L78, T80, L81, T87, G88, Y89, H90, G91, T94, R95, D96, F97, Y167, F168, H169, H170, N171, G172, D173, I174, S175, N176, D178, D179, R180, Y181, E182, A183, Q184, K186, N187, F188, T189.

In more preferred embodiment, the variant of a maltogenic alpha-amylase comprises a substitution corresponding to one or more of the following substitutions in the amino acid sequence set forth in SEQ ID NO: 1:

D17E/Q, A30M/L/A/V/I/E/Q, S32D/E/N/Q, R95M/L/A/V/I/E/Q, H103Y/N/Q/D/E, N131D, Q201E, I174E/Q, and H169N/D/E/Q

V74I, L75N/D/Q/I/V, L78N/I, T80I/LV/S/N/G, L81I/V/S/T/N/Q/K/H, T87S/I, G88A/S/T, Y89F, H90N/Q/K, G91A/S/T, T94N/D/A/M/V/I, R95K/Q, D96N/V/Q/I, F97Y, Y167F/R/C, F168Y, H169N/Q/K, H170N/Q/K, N171D/E/Q/H/R/K/G, G172A/T/S, D173N/S/T/Y/R/G, I174N/Q/L, S175T/A/N/D, N176S/T/H/Q/P, D178N/Q/E/K/H, D179Y/N/H, R180W, Y181R/F/C/L, E182D, A183S/C/G, Q184E, K186R, N187Q/E/L/F/H/K/V/L, F188Y/L/I/H/N, T189N/D/A/S/H/Y/G.

In another preferred embodiment of the invention with respect to altering the 35 Ca<sup>2+</sup> binding of a maltogenic alpha-amylase the partial sequence N28-P29-A30-K31-S32-Y33-G34 as set forth in SEQ ID NO: 1 is modified.

Similar substitutions may be introduced in equivalent positions of other maltogenic alpha-amylases. Modifications of particular interest are any combination of one or more of the above with any of the other modifications disclosed herein.

### Other substitutions

Variants with improved stability of the enzyme can be achieved by improving existing or introducing new interdomain and intradomain contacts. Such improved stability can be achieved by the modifications listed below.

The maltogenic alpha-amylase having the amino acid sequence shown in SEQ ID NO: 1 may be stabilized by the introduction of one or more interdomain disulfide bonds. Accordingly, another preferred embodiment of the present invention relates to a variant of a parent maltogenic alpha-amylase which has improved stability and at least one more interdomain disulfide bridge as compared to said parent, wherein said variant comprises a modification in a position corresponding to at least one of the following pairs of positions in SEQ ID NO: 1:

G236 + S583, G618 + R272, T252 + V433 and/or A348 + V487.

In a more preferred embodiment, the substitution corresponds to at least one of the following pairs:

G236C + S583C, G618C + R272C, T252C + V433C and/or A348C + V487C.

Another preferred embodiment of the invention relates to a variant of a parent maltogenic alpha-amylase which has an improved stability and an altered interdomain interaction as compared to said parent, wherein said variant comprises a substitution in a position corresponding at least one of the following sets of positions in SEQ ID NO: 1:

- i) F143, F194, L78;
- ii) A341, A348, L398, I415, T439, L464, L465;
- 25 iii) L557;

15

35

- iv) S240, L268;
- v) Q208, L628;
- vi) F427, Q500, N507, M508, S573; and
- vii) 1510, V620.

In a more preferred embodiment, the substitution corresponds to at least one of the following sets:

- i) F143Y, F194Y, L78Y/F/W/E/Q;
- ii) A341S/D/N, A348V/I/L, L398E/Q/N/D, I415E/Q, T439D/E/Q/N, L464D/E, L465D/E/N/Q/R/K;
  - iii) L557Q/E/N/D;
  - iv) S240D/E/N/Q, L268D/E/N/Q/R/K;
  - v) Q208D/E/Q, L628E/Q/N/D;

vi) F427E/Q/R/K/Y, Q500Y, N507Q/E/D, M508K/R/E/Q, S573D/E/N/Q; and/or vii) I510D/E/N/Q/S, V620D/E/N/Q.

Another preferred embodiment of the invention relates to a variant of a parent maltogenic alpha-amylase which has an improved stability and one or more salt bridges as compared to said parent, wherein said variant comprises a substitution in a position corresponding at least one of the following sets of positions in SEQ ID NO: 1:

N106, N320 and Q624.

In a more preferred embodiment, the variant of a maltogenic alpha-amylase comprises a substitution corresponding to the following substitutions in the amino acid sequence set forth in SEQ ID NO: 1:

N106R, N320E/D and/or Q624E.

Another embodiment of the invention relates to a variant of a parent maltogenic alpha-amylase which has an improved stability and wherein said variant comprises a substitution in a position corresponding at least one of the following sets of positions in SEQ ID NO: 1:

K40, V74, S141, T142, F188, N234, K249, D261, D261, L268, V279, N342, G397, A403, K425, S442, S479, S493, T494, S495, A496, S497, A498, Q500, K520, A555 and N595.

In a more preferred embodiment, the variant of a maltogenic alpha-amylase comprises a substitution corresponding to one or more of the following substitutions with proline in the amino acid sequence set forth in SEQ ID NO: 1:

V74P, S141P, N234P, K249P, L268P, V279P, N342P, G397P, A403P, S442P, S479P, S493P, T494P, S495P, A496P, S497P, A498P, Q500P, and/or A555P.

Other preferred substitutions are K40R, T142A, F188I/L, D261G, K425E, 25 K520R, and/or N595I.

Analogously, it may be preferred that one or more histidine residues present in the parent maltogenic alpha-amylase is or are substituted with a non-histidine residues such as Y, V I, L, F, M, E, Q, N, or D. Accordingly, in another preferred embodiment, the variant of a maltogenic alpha-amylase comprises a substitution of an amino acid residue corresponding to one or more of the following residues of the amino acid sequence set forth in SEQ ID NO: 1:

H103, H220, and H344

In a more preferred embodiment, the variant of a maltogenic alpha-amylase comprises a substitution corresponding to one or more of the following substitutions in the amino acid sequence set forth in SEQ ID NO: 1:

H103Y/V/I/L/F/Y, H220Y/L/M, and H344E/Q/N/D/Y.

It may be preferred that one or more asparagine or glutamine residues present in the parent maltogenic alpha-amylase is or are substituted with a residue lacking the

30

35

amide on the side chain. Accordingly, in another preferred embodiment, the variant of a Novamyl-like comprises a substitution of an amino acid residue corresponding to one or more of the following residues of the amino acid sequence set forth in SEQ ID NO: 1:

Q13, N26, N77, N86, N99, Q119, N120, N131, N152, N171, N176, N187, S Q201, N203, N234, Q247, N266, N275, N276, N280, N287, Q299, N320, N327, N342, Q365, N371, N375, N401, N436, N454, N468, N474, Q500, N507, N513, Q526, N575, Q581, N621, Q624 and N664.

In more preferred embodiment, the variant of a maltogenic alpha-amylase comprises a substitution corresponding to one or more of the following substitutions in the amino acid sequence set forth in SEQ ID NO: 1:

Q13S/T/A/V/L/I/F/M, N26S/T/A/V/L/I, N77S/T/A/V/L/I, N86S/T/A/V/L/I, N99T/S/V/L, Q119T/S, N120S/T/A/V/L/I, N131S/T/A/V/L/I, N152T/S/V/L, N171Y/D/S/T, N187S/T/A/V/L/I, Q201S/T/A/V/L/I/F/M, N203D/S/T/A/V/L/I, N176S/T/A/V/L/I, Q247S/T/AV/L/I/F/M, N266S/T/A/V/L/I. N275S/T/A/V/L/I, N234S/T/A/V/L/I. 15 N276S/T/AV/L/I, N280S/T/AV/L/I, N287S/T/AV/L/I, Q299L/T/S, N320S/T/AV/L/I, N327S/T/AV/L/I, N342S/T/AV/L/I, Q365S/T/AV/L/I, N371S/T/AV/L/I, N375S/T/AV/L/I, N436S/T/A/V/L/I, N454D/S/T/A/V/L/I, N468D/S/T/A/V/L/I, N401S/T/A/V/L/I, N474D/S/T/AV/L/I, Q500S/T/AV/L/I/F/M, N507S/T/AV/L/I, N513S/T/AV/L/I, Q526 D/S/T/A/V/L/I. N575S/T/A/V/L/I. Q581S/T/AV/L/I/F/M, N621S/T/A/V/L/I 20 Q624S/T/A/V/L/I/F/M and N664D/S/T/A/V/L/I.

Another embodiment of the invention relates to a variant of a parent maltogenic alpha-amylase which has improved stability and improved hydrogen bond contacts as compared to said parent, wherein said variant comprises a modification in a position corresponding to one or more of the following positions in SEQ ID NO: 1:

I16, L35, M45, P73, D76, D79, A192, I100, A148, A163+G172, L268, V281, D285, L321, F297, N305, K316, S573, A341, M378, A381, F389, A483, A486, I510, A564, F586, K589, F636, K645, A629, and/or T681.

In a preferred embodiment, the modification corresponds to one or more of the following:

I16T/D/N, L35Q, M45K, P73Q, D76E, D79E/Y, A192S/D/N, I100T/S/D/N/E/Q, A148D/N/E/Q/S/T/R/K, A163Y+G172S/D/N, L268R/K, V281/Q, D285R/K, L321Q, F297N/D/Q/E, N305K/R, K316N/D, S573N/D, A341R/K, M378R/K, A381S/D/N, F389Y, A483S/D/N, A486Q/E, I510R/K, A564S/D/N, F586S/D/N, K589S/D/Q/N, F636Y, K645T, A629N/D/E/Q, and/or T681D/N/E/Q/S.

Similar substitutions may be introduced in equivalent positions of other maltogenic alpha-amylases. Substitutions of particular interest are any combination of one or more of the above with any of the other modifications disclosed herein.

Before actually constructing a maltogenic alpha-amylase variant to achieve any of the above objectives, it may be convenient to evaluate whether or not the contemplated amino acid modification can be accommodated into the maltogenic alpha-amylase structure, e.g. into a model of the three-dimensional structure of the parent maltogenic alpha-amylase.

# Maltogenic alpha-amylase variants with altered thermostability and/or altered temperature dependent activity profile

The invention further relates to a variant of a parent maltogenic alpha-amylase, which results from substitution, deletion or insertion of one or more amino acid residues so as to obtain a variant having an altered thermostability or temperature dependent activity profile.

The structure of the maltogenic alpha-amylase contains a number of unique internal cavities which may contain water and a number of crevices. In order to increase the thermostability of the polypeptide it may be desirable to reduce the number or size of cavities and crevices, e.g., by introducing one or more hydrophobic contacts, preferably achieved by introducing amino acids with bulkier side groups in the vicinity or surroundings of the cavity. For instance, the amino acid residues to be modified are those which are involved in the formation of the cavity.

Accordingly, in a further aspect the present invention relates to a method of increasing the thermostability and/or altering the temperature dependent activity profile of a parent maltogenic alpha-amylase, which method comprises:

- i) identifying an internal cavity or a crevice of the parent maltogenic alphaamylase in the three-dimensional structure of said polypeptide;
- ii) substituting, in the structure, one or more amino acid residues in the neighbourhood of the cavity or crevice identified in step i) with another amino acid residue which, from structural or functional considerations, is determined to increase the hydrophobic interaction and to fill out or reduce the size of the cavity or crevice; and
- iii) constructing a variant of the parent maltogenic alpha-amylase resulting from step ii) and testing the thermostability and/or temperature dependent activity of the variant.

The structure identified in Appendix 1 may be used for identifying the cavity or crevice of the parent maltogenic alpha-amylase.

It will be understood that the cavity or crevice is identified by the amino acid residues surrounding said cavity or crevice, and that modification of said amino acid residues are of importance for filling or reducing the size of said cavity or crevice. Preferably, the modification is a substitution with a bulkier amino acid residue, i.e. one with a greater side chain volume. For example, all the amino acids are bulkier than Gly,

whereas Tyr and Trp are bulkier than Phe. The particular amino acid residues referred to below are those which in a crystal structure have been found to flank the cavity or crevice in question.

In a preferred embodiment, the variant of a maltogenic alpha-amylase, in order to fill, either completely or partly, cavities located internally in the structure, comprises a substitution of an amino acid residue corresponding to one or more of the following residues of the amino acid sequence set forth in SEQ ID NO: 1:

L51, L75, L78, G88, G91, T94, V114, I125, V126, T134, G157, L217, S235, G236, V254, V279, V281, L286, V289, I290, V308, L321, I325, D326, L343, F349, S353, I359, I405, L448, Q449, L452, I470, G509, V515, S583, G625, L627, L628 and A670.

L71, S72, V74, L75, L78, T80, L81, G83, T84, D85, N86, T87, G88, Y89, H90, G91, T94, R95, D96, F97, Y167, F168, H169, H170, N171, G172, D173, I174, S175, N176, D178, D179, R180, Y181, E182, A183, Q184, K186, N187, F188, T189, D190, A192, G193, F194, S195, L196.

In a more preferred embodiment, the variant of a maltogenic alpha-amylase comprises one or more substitutions corresponding to the following substitutions in the amino acid sequence set forth in SEQ ID NO: 1:

L217 in combination with L75 (e.g. L217F/Y in combination with L75F/Y), 20 L51W, L75F/Y, L78I, G88AV/T, G91T/S/V/N, T94V/I/L, V114V/I/L, I125L/M/F/Y/W, V126I/L, T134V/I/L/M/F/Y/W, G157AV/I/L, L217V/I/M/F/Y/W, S235I/L/M/F/Y/W, G236AV/I/L/M/F/Y/W, V254I/L/M/F/Y/W, V279M/I/L/F, V281I/L/M/F/Y/W, L286F, V289I/L/R, I290M/L/F, V308I/L/M/F/Y/W, L321I/M/F/Y/W, I325L/M/F/Y/W, D326E/Q, L343M/F/Y/W, F349W/Y, S353V/I/L, I359L/M/F/Y/W, I405M/L/Y/F/W, L448Y, Q449Y, L452M/Y/F/W, I470M/L/F, G509AV/I/L/M/S/T/D/N, V515I/L, S583V/I/L/V, G625AV/I/L/M/F/Y/W, L627M/F/Y, L628M/I/F/Y/W and A670V/I/L/M/F/Y/W,

L71I, S72C, V74I, L75N/D/Q/I/V, L78N/I, T80I/L/V/S/N/G, L81I/V/S/T/N/Q/K/H, G83A/S/T/N/Q/E/D/R/H/L, T84S/A/N/D/G, D85A/T/S/N/G, N86Q/E/D/Y/H/K, T87S/I, G88A/S/T, Y89F, H90N/Q/K, G91A/S/T, T94N/D/A/M/V/I, R95K/Q, D96N/V/Q/I, F97Y, Y167F/R/C, F168Y, H169N/Q/K, H170N/Q/K, N171D/E/Q/H/R/K/G, G172A/T/S, D173N/S/T/Y/R/G, I174N/Q/L, S175T/A/N/D, N176S/T/H/Q/P, D178N/Q/E/K/H, D179Y/N/H, R180W, Y181R/F/C/L, E182D, A183S/C/G, Q184E, K186R, N187Q/E/L/F/H/K/V/L, F188Y/L/I/H/N, T189N/D/A/S/H/Y/G, D190E/Q/H/N/K, A192T/D/E/N/K, G193A/S/T, F194Y, S195N/D/E/R/K/G, L196I.

Similar substitutions may be introduced in equivalent positions of other maltogenic alpha-amylases. Variants of particular interest have a combination of one or more of the above with any of the other modifications disclosed herein.

### Maltogenic alpha-amylase variants with an alter d cleavag pattern

One aim of the present invention is to change the degradation characteristics of a maltogenic alpha-amylase. Thus, Novamyl hydrolyzes starch to form predominantly maltose (G2) and a small amount of glucose (G1), but virtually no higher oligosaccharides (G3+). It may be desirable to change this cleavage pattern, e.g. so as to form higher amounts of higher oligosaccharides, such as maltotriose (G3), maltotetraose (G4) and maltopentaose (G5).

A variant of a parent maltogenic alpha-amylase in which the substrate cleavage pattern is altered as compared to said parent may be constructed by a method which to comprises:

- i) identifying the substrate binding area of the parent maltogenic alpha-amylase in a model of the three-dimensional structure, e.g. within a sphere of 4 Å from the substrate binding site as defined in the section above entitled "Substrate Binding Site";
- ii) substituting in the model one or more amino acid residues of the substrate binding area of the cleft identified in i) which is or are believed to be responsible for the cleavage pattern of the parent with another amino acid residue which from structural or functional considerations is believed to result in an altered substrate cleavage pattern, or deleting one or more amino acid residues of the substrate binding area contemplated to introduce favorable interactions to the substrate or adding one or more amino acid residues to the substrate binding area contemplated to introduce favorable interactions to the substrate; and
  - iii) constructing a maltogenic alpha-amylase variant resulting from step ii) and testing the substrate cleavage pattern of the variant.

Accordingly, another aspect of the invention relates to a variant of a parent maltogenic alpha-amylase which has an altered substrate binding site as compared to said parent, which variant comprises a modification in a position corresponding to one or both of the following positions in SEQ ID NO: 1:

V281 and/or A629.

In a preferred embodiment, the variant comprises a modification corresponding to:

V281Q and/or A629N/D/E/Q.

Similar modifications may be introduced in equivalent positions of other maltogenic alpha-amylases. Substitutions of particular interest are any combination of one or both of the above with any of the other modifications disclosed herein.

# Maltogenic alpha-amylase variants with improved ability to reduce retrogradation of starch and/or staling of bread

The invention provides maltogenic alpha-amylase variants having improved ability to reduce the retrogradation of starch and/or the staling of bread. Preferred variants comprise a modification at one or more positions corresponding to the following amino acid residues in SEQ ID NO: 1:

A30, K40, N115, T142, F188, T189, P191, A192, G193, F194, S195, D261, N327, K425, K520 and N595.

In a more preferred embodiment, the variant comprises one or more modifications corresponding to the following in SEQ ID NO: 1:

A30D, K40R, N115D, T142A, F188L, T189Y,  $\Delta$  (191-195), D261G, D261G, N327S, K425E, K520R and N595I.

# Determination of residues within 10Å from calcium ions

The coordinates of Appendix 1 are read into the INSIGHT program (BIOSYM Technologies). The spatial coordinates are presented showing the bonds between the atoms. The ions are presented as well as the water atoms. The part of the program package for creating subsets was used to create a 10 Å subset around the calcium ions in the structure by using the command ZONE. All residues identified as having an atom within the designated 10 Å distance from a calcium ion are compiled and listed by using the command LIST MOLECULE. By giving the ions the name "VAT CA" in the coordinate file, a 10 Å sphere around all atoms called "VAT CA" is compiled. The specific residues identified in this manner are given further above in the section entitled "Calcium binding".

### 25 Determination of cavities

The solved structure of Novamyl with the structural coordinates set forth in Appendix 1 reveals many internal crevices and cavities. When analysing for such cavities the Connolly program is normally used (Lee, B. and Richards, F.M. (1971) J. Mol. Biol. 55:379-400). The program uses a probe with radius to search the external and internal surface of the protein. The smallest crevice observable in this way has the probe radius.

To analyse the solved structure a modified version of the Connolly program included in the program of INSIGHT was used. In the first step, the water molecules and the ions were removed by unmerging these atoms from the solved structure. By using the command MOLECULE SURFACE SOLVENT the solvent accessible surface area was calculated for all atoms and residues using a probe radius of 1.4 Å, and

25

displayed graphically together with the model of the solved structure. The internal cavities are then seen as dot surfaces with no connections to the external surface.

Suggestions for specific modifications to fill out the cavities are given above in the section entitled "Variants with altered thermostability and/or altered temperature 5 dependent activity profile"). By using the homology built structures or/and comparisons based on sequence alignment, mutations for homologous structures of maltogenic alpha-amylases can be made.

### Nomenclature for amino acid modifications

The nomenclature used herein for defining mutations is essentially as described in WO 92/05249. Thus, F188H indicates a substitution of the amino acid F (Phe) in position 188 with the amino acid H (His). V129S/T/G/V indicates a substitution of V129 with S, T, G or V.  $\Delta$  (191-195) or  $\Delta$  (191-195) indicates a deletion of amino acids in positions 191-195. 192-A-193 indicates an insertion of A between amino acids 15 192 and 193.

### Polypeptide sequence identity

For purposes of the present invention, the degree of identity may be suitably determined according to the method described in Needleman, S.B. and Wunsch, C.D., (1970), Journal of Molecular Biology, 48, 443-45, with the following settings for 20 polypeptide sequence comparison: GAP creation penalty of 3.0 and GAP extension penalty of 0.1. The determination may be done by means of a computer program known such as GAP provided in the GCG program package (Program Manual for the Wisconsin Package, Version 8, August 1994, Genetics Computer Group, 575 Science Drive, Madison, Wisconsin, USA 53711).

The variants of the invention have an amino acid identity with amino acids 1-686 of SEQ ID NO: 1 of at least 70 %, preferably at least 80 %, e.g. at least 90 %, particularly at least 95 % or at least 98 %.

### Hybridization

Suitable experimental conditions for determining hybridization between a 30 nucleotide probe and a homologous DNA or RNA sequence involves presoaking of the filter containing the DNA fragments or RNA to hybridize in 5x SSC (sodium chloride/sodium citrate, Sambrook, et al., 1989) for 10 min, and prehybridization of the filter in a solution of 5x SSC, 5x Denhardt's solution (Sambrook, et al., 1989), 0.5% SDS and 100 µg/ml of denatured sonicated salmon sperm DNA (Sambrook, et al., 35 1989), followed by hybridization in the same solution containing a random-primed

30

(Feinberg, A. P. and Vogelstein, B. (1983) Anal. Biochem. 132:6-13), 32P-dCTP-labeled (specific activity > 1 x 109 cpm/µg ) probe for 12 hours at ca. 45°C. The filter is then washed twice for 30 minutes in 2x SSC, 0.5% SDS at least 55°C (low stringency), preferably at least 60°C (medium stringency), more preferably at least 65°C 5 (medium/high stringency), more preferably at least 70°C (high stringency), even more preferably at least 75°C (very high stringency).

Molecules which hybridize to the oligonucleotide probe under these conditions are detected by exposure to x-ray film.

# Methods of preparing variants of maltogenic alpha-amylases

### 10 Cloning a DNA sequence encoding a Novamyl-like polypeptide

The DNA sequence encoding a parent maltogenic alpha-amylase may be isolated from any cell or microorganism producing the maltogenic alpha-amylase in question, using various methods well known in the art, for example, from the Bacillus strain NCIB 11837.

First, a genomic DNA and/or cDNA library should be constructed using chromosomal DNA or messenger RNA from the organism that produces the maltogenic alpha-amylase to be studied. Then, if the amino acid sequence of the  $\alpha$ -amylase is known, homologous, labelled oligonucleotide probes may be synthesised and used to identify maltogenic alpha-amylase-encoding clones from a genomic library prepared 20 from the organism in question. Alternatively, a labelled oligonucleotide probe containing sequences homologous to a known α-amylase gene could be used as a probe to identify maltogenic alpha-amylase-encoding clones, using hybridization and washing conditions of lower stringency.

Another method for identifying maltogenic alpha-amylase-encoding clones 25 involves inserting fragments of genomic DNA into an expression vector, such as a plasmid, transforming α-amylase negative bacteria with the resulting genomic DNA library, and then plating the transformed bacteria onto agar containing a substrate for maltogenic alpha-amylase, thereby allowing clones expressing maltogenic alphaamylase activity to be identified.

Alternatively, the DNA sequence encoding the enzyme may be prepared synthetically by established standard methods, e.g. the phosphoroamidite method described by S.L. Beaucage and M.H. Caruthers (1981) or the method described by Matthes et al. (1984). In the phosphoroamidite method, oligonucleotides are synthesized, e.g. in an automatic DNA synthesizer, purified, annealed, ligated and cloned 35 in appropriate vectors.



Finally, the DNA sequence may be of mixed genomic and synthetic origin, mixed synthetic and cDNA origin or mixed genomic and cDNA origin, prepared by ligating fragments of synthetic, genomic or cDNA origin, wherein the fragments correspond to various parts of the entire DNA sequence, in accordance with techniques well known in the art. The DNA sequence may also be prepared by polymerase chain reaction (PCR) using specific primers, for instance as described in US 4,683,202 or R.K. Saiki et al. (1988).

### Site-directed Mutagenesis

Once a maltogenic alpha-amylase-encoding DNA sequence has been isolated, and desirable sites for modification identified, modifications may be introduced using synthetic oligonucleotides. These oligonucleotides contain nucleotide sequences flanking the desired modification sites; mutant nucleotides are inserted during oligonucleotide synthesis. In a specific method, a single-stranded gap of DNA, bridging the maltogenic alpha-amylase-encoding sequence, is created in a vector carrying the maltogenic alpha-amylase gene. Then the synthetic nucleotide, bearing the desired modification, is annealed to a homologous portion of the single-stranded DNA. The remaining gap is then filled in with DNA polymerase I (Klenow fragment) and the construct is ligated using T4 ligase. A specific example of this method is described in Morinaga et al. (1984): US 4,760,025 discloses the introduction of oligonucleotides encoding multiple modifications by performing minor alterations of the cassette. However, an even greater variety of modifications can be introduced at any one time by the Morinaga method because a multitude of oligonucleotides, of various lengths, can be introduced.

Another method of introducing modifications into a maltogenic alpha-amylase25 encoding DNA sequences is described in Nelson and Long (1989). It involves a 3-step
generation of a PCR fragment containing the desired modification introduced by using
a chemically synthesized DNA strand as one of the primers in the PCR reactions. From
the PCR-generated fragment, a DNA fragment carrying the modification may be
isolated by cleavage with restriction endonucleases and reinserted into an expression
plasmid.

### Random Mutagenesis

35

Random mutagenesis is suitably performed either as localised or regionspecific random mutagenesis in at least three parts of the gene translating to the amino acid sequence shown in question, or within the whole gene.

The random mutagenesis of a DNA sequence encoding a parent maltogenic alpha-amylase may be conveniently performed by use of any method known in the art.



In relation to the above, a further aspect of the present invention relates to a method for generating a variant of a parent Novamyl-like  $\alpha$ -amylase, wherein the variant exhibits increased stability at low pH and at low calcium concentration relative to the parent, the method comprising:

- (a) subjecting a DNA sequence encoding the parent Novamyl-like  $\alpha$ -amylase to random mutagenesis,
- (b) expressing the mutated DNA sequence obtained in step (a) in a host cell, and
- (c) screening for host cells expressing a Novamyl-like  $\alpha$ -amylase variant which has an altered property relative to the parent Novamyl-like  $\alpha$ -amylase.
  - Step (a) of the above method of the invention is preferably performed using doped primers, as described in the working examples herein (*vide infra*).

For instance, the random mutagenesis may be performed by use of a suitable physical or chemical mutagenizing agent, by use of a suitable oligonucleotide, or by subjecting the DNA sequence to PCR generated mutagenesis. Furthermore, the random mutagenesis may be performed by use of any combination of these mutagenizing agents. The mutagenizing agent may, e.g., be one which induces transitions, transversions, inversions, scrambling, deletions, and/or insertions.

Examples of a physical or chemical mutagenizing agent suitable for the present purpose include ultraviolet (UV) irradiation, hydroxylamine, N-methyl-N'-nitro-N-nitrosoguanidine (MNNG), O-methyl hydroxylamine, nitrous acid, ethyl methane sulphonate (EMS), sodium bisulphite, formic acid, and nucleotide analogues. When such agents are used, the mutagenesis is typically performed by incubating the DNA sequence encoding the parent enzyme to be mutagenized in the presence of the mutagenizing agent of choice under suitable conditions for the mutagenesis to take place, and selecting for mutated DNA having the desired properties.

When the mutagenesis is performed by the use of an oligonucleotide, the oligonucleotide may be doped or spiked with the three non-parent nucleotides during the synthesis of the oligonucleotide at the positions which are to be changed. The doping or spiking may be done so that codons for unwanted amino acids are avoided. The doped or spiked oligonucleotide can be incorporated into the DNA encoding the maltogenic alpha-amylase enzyme by any published technique, using e.g. PCR, LCR or any DNA polymerase and ligase as deemed appropriate.

Preferably, the doping is carried out using "constant random doping", in which the percentage of wild-type and modification in each position is predefined. Furthermore, the doping may be directed toward a preference for the introduction of certain nucleotides, and thereby a preference for the introduction of one or more specific amino acid residues. The doping may be made, e.g., so as to allow for the

15

introduction of 90% wild type and 10% modifications in each position. An additional consideration in the choice of a doping scheme is based on genetic as well as proteinstructural constraints. The doping scheme may be made by using the DOPE program which, inter alia, ensures that introduction of stop codons is avoided.

When PCR-generated mutagenesis is used, either a chemically treated or nontreated gene encoding a parent maltogenic alpha-amylase enzyme is subjected to PCR under conditions that increase the misincorporation of nucleotides (Deshler 1992; Leung et al., Technique, Vol.1, 1989, pp. 11-15).

A mutator strain of E. coli (Fowler et al., Molec. Gen. Genet., 133, 1974, pp. 10 179-191), S. cereviseae or any other microbial organism may be used for the random mutagenesis of the DNA encoding the maltogenic alpha-amylase by, e.g., transforming a plasmid containing the parent enzyme into the mutator strain, growing the mutator strain with the plasmid and isolating the mutated plasmid from the mutator strain. The mutated plasmid may be subsequently transformed into the expression organism.

The DNA sequence to be mutagenized may be conveniently present in a genomic or cDNA library prepared from an organism expressing the parent maltogenic alpha-amylase. Alternatively, the DNA sequence may be present on a suitable vector such as a plasmid or a bacteriophage, which as such may be incubated with or otherwise exposed to the mutagenising agent. The DNA to be mutagenized may also be 20 present in a host cell either by being integrated in the genome of said cell or by being present on a vector harbored in the cell. Finally, the DNA to be mutagenized may be in isolated form. It will be understood that the DNA sequence to be subjected to random mutagenesis is preferably a cDNA or a genomic DNA sequence.

In some cases it may be convenient to amplify the mutated DNA sequence 25 prior to performing the expression step b) or the screening step c). Such amplification may be performed in accordance with methods known in the art, the presently preferred method being PCR-generated amplification using oligonucleotide primers prepared on the basis of the DNA or amino acid sequence of the parent enzyme.

Subsequent to the incubation with or exposure to the mutagenising agent, the 30 mutated DNA is expressed by culturing a suitable host cell carrying the DNA sequence under conditions allowing expression to take place. The host cell used for this purpose may be one which has been transformed with the mutated DNA sequence, optionally present on a vector, or one which was carried the DNA sequence encoding the parent enzyme during the mutagenesis treatment. Examples of suitable host cells are the 35 following: gram positive bacteria such as Bacillus subtilis, Bacillus licheniformis, Bacillus lentus, Bacillus brevis, Bacillus stearothermophilus, Bacillus alkalophilus, Bacillus amyloliquefaciens, Bacillus coagulans, Bacillus circulans, Bacillus lautus, Bacillus

megaterium, Bacillus thuringiensis, Streptomyces lividans or Streptomyces murinus; and gram negative bacteria such as E. coli.

The mutated DNA sequence may further comprise a DNA sequence encoding functions permitting expression of the mutated DNA sequence.

### 5 Localized random mutagenesis

The random mutagenesis may be advantageously localised to a part of the parent maltogenic alpha-amylase in question. This may, e.g., be advantageous when certain regions of the enzyme have been identified to be of particular importance for a given property of the enzyme, and when modified are expected to result in a variant having improved properties. Such regions may normally be identified when the tertiary structure of the parent enzyme has been elucidated and related to the function of the enzyme.

The localized, or region-specific, random mutagenesis is conveniently performed by use of PCR generated mutagenesis techniques as described above or any other suitable technique known in the art. Alternatively, the DNA sequence encoding the part of the DNA sequence to be modified may be isolated, e.g., by insertion into a suitable vector, and said part may be subsequently subjected to mutagenesis by use of any of the mutagenesis methods discussed above.

For region-specific random mutagenesis with a view to improving the stability of calcium binding of a parent maltogenic alpha-amylase, codon positions corresponding to the following amino acid residues from the amino acid sequence set forth in SEQ ID NO: 1 may appropriately be targeted:

Residues:Regions:

16-33, 35-36, 40: 16-40

25 46-54, 56: 46-56

73-81: 73-81

87-89, 91, 93-96, 99-105, 109: 87-109

129-134, (145, 150): 129-134

167-172, 174, 177, 180-189: 167-189

196-202, 206-210: 196-210

228-235, 237: 228-237

378

30

637

With a view to achieving improved binding of a substrate, i.e., improved binding of a carbohydrate species, such as amylose or amylopectin, by a maltogenic alphaamylase variant with a modified, e.g. higher, substrate specificity and/or a modified, e.g. higher, specificity with respect to cleavage, i.e. hydrolysis, of the substrate, it appears

that the following codon positions in the following regions of the amino acid sequence shown in SEQ ID NO: 1, may particularly appropriately be targeted for modification by region-specific mutagenesis:

70-97, 127-143, 174-198, 226-233, 255-270, 282-292, 324-331, 370-376.

For region-specific random mutagenesis with a view to altering the substrate specificity and/or the pH dependent activity profile, the following regions of SEQ ID NO: 1 may be targeted: 70-97, 174-198.

The following regions may be targeted with a view to improving the thermostability: 70-109, 167-200.

# 10 General method for random mutagenesis by use of the DOPE program

The random mutagenesis may be carried out by the following steps:

- 1. Select regions of interest for modification in the parent enzyme
- 2.Decide on mutation sites and nonmutated sites in the selected region
- 3.Decide on which kind of mutations should be carried out, e.g. with respect to the desired stability and/or performance of the variant to be constructed
  - 4. Select structurally reasonable mutations
  - 5. Adjust the residues selected by step 3 with regard to step 4.
  - 6. Analyse by use of a suitable dope algorithm the nucleotide distribution.
- 7.If necessary, adjust the wanted residues to genetic code realism, e.g. taking into account constraints resulting from the genetic code, e.g. in order to avoid introduction of stop codons; the skilled person will be aware that some codon combinations cannot be used in practice and will need to be adapted
  - 8.Make primers
  - 9.Perform random mutagenesis by use of the primers
- 10. Select resulting  $\alpha$ -amylase variants by screening for the desired improved properties.

Suitable dope algorithms for use in step 6 are well known in the art. One such algorithm is described by Tomandl, D. et al., 1997, Journal of Computer-Aided Molecular Design 11:29-38. Another algorithm is DOPE (Jensen, LJ, Andersen, KV, 30 Svendsen, A, and Kretzschmar, T (1998) Nucleic Acids Research 26:697-702).

### Expression of maltogenic alpha-amylase variants

The construction of the variant of interest is accomplished by cultivating a microorganism comprising a DNA sequence encoding the variant under conditions which are conducive for producing the variant, and optionally subsequently recovering the variant from the resulting culture broth. This is described in detail further below.

According to the invention, a DNA sequence encoding the variant produced by methods described above, or by any alternative methods known in the art, can be expressed, in the form of a protein or polypeptide, using an expression vector which typically includes control sequences encoding a promoter, operator, ribosome binding site, translation initiation signal, and, optionally, a repressor gene or various activator genes.

The recombinant expression vector carrying the DNA sequence encoding an maltogenic alpha-amylase variant of the invention may be any vector which may conveniently be subjected to recombinant DNA procedures, and the choice of vector will often depend on the host cell into which it is to be introduced. Thus, the vector may be an autonomously replicating vector, i.e. a vector which exists as an extrachromosomal entity, the replication of which is independent of chromosomal replication, e.g. a plasmid, a bacteriophage or an extrachromosomal element, minichromosome or an artificial chromosome. Alternatively, the vector may be one which, when introduced into a host cell, is integrated into the host cell genome and replicated together with the chromosome(s) into which it has been integrated.

In the vector, the DNA sequence should be operably connected to a suitable promoter sequence. The promoter may be any DNA sequence which shows transcriptional activity in the host cell of choice and may be derived from genes encoding proteins either homologous or heterologous to the host cell. Examples of suitable promoters for directing the transcription of the DNA sequence encoding a maltogenic alpha-amylase variant of the invention, especially in a bacterial host, are the promoter of the *lac* operon of *E.coli*, the *Streptomyces coelicolor* agarase gene *dag*A promoters, the promoters of the *Bacillus licheniformis* α-amylase gene (*amyL*), the promoters of the *Bacillus stearothermophilus* maltogenic amylase gene (*amyM*), the promoters of the *Bacillus amyloliquefaciens* α-amylase (*amyQ*), the promoters of the *Bacillus subtilis* xylA and xylB genes, etc. For transcription in a fungal host, examples of useful promoters are those derived from the gene encoding *A. oryzae* TAKA amylase, *Rhizomucor miehei* aspartic proteinase, *A. niger* neutral α-amylase, *A. niger* acid stable α-amylase, *A. niger* glucoamylase, *Rhizomucor miehei* lipase, *A. oryzae* alkaline protease, *A. oryzae* triose phosphate isomerase or *A. nidulans* acetamidase.

The expression vector of the invention may also comprise a suitable transcription terminator and, in eukaryotes, polyadenylation sequences operably connected to the DNA sequence encoding the maltogenic alpha-amylase variant of the invention. Termination and polyadenylation sequences may suitably be derived from the same sources as the promoter.

The vector may further comprise a DNA sequence enabling the vector to replicate in the host cell in question. Examples of such sequences are the origins of replication of plasmids pUC19, pACYC177, pUB110, pE194, pAMB1 and pIJ702.

The vector may also comprise a selectable marker, e.g. a gene the product of which complements a defect in the host cell, such as the dal genes from B. subtilis or B. licheniformis, or one which confers antibiotic resistance such as ampicillin, kanamycin, chloramphenicol or tetracycline resistance. Furthermore, the vector may comprise Aspergillus selection markers such as amdS, argB, niaD and sC, a marker giving rise to hygromycin resistance, or the selection may be accomplished by co-transformation, e.g. as described in WO 91/17243.

While intracellular expression may be advantageous in some respects, e.g. when using certain bacteria as host cells, it is generally preferred that the expression is extracellular. In general, the *Bacillus* α-amylases mentioned herein comprise a preregion permitting secretion of the expressed protease into the culture medium. If desirable, this preregion may be replaced by a different preregion or signal sequence, conveniently accomplished by substitution of the DNA sequences encoding the respective preregions.

The procedures used to ligate the DNA construct of the invention encoding maltogenic alpha-amylase variant, the promoter, terminator and other elements, respectively, and to insert them into suitable vectors containing the information necessary for replication, are well known to persons skilled in the art (cf., for instance, Sambrook et al. (1989)).

The cell of the invention, either comprising a DNA construct or an expression vector of the invention as defined above, is advantageously used as a host cell in the recombinant production of a maltogenic alpha-amylase variant of the invention. The cell may be transformed with the DNA construct of the invention encoding the variant, conveniently by integrating the DNA construct (in one or more copies) in the host chromosome. This integration is generally considered to be an advantage as the DNA sequence is more likely to be stably maintained in the cell. Integration of the DNA constructs into the host chromosome may be performed according to conventional methods, e.g. by homologous or heterologous recombination. Alternatively, the cell may be transformed with an expression vector as described above in connection with the different types of host cells.

The cell of the invention may be a cell of a higher organism such as a mammal or an insect, but is preferably a microbial cell, e.g. a bacterial or a fungal (including yeast) cell.

Examples of suitable bacteria are gram positive bacteria such as Bacillus subtilis, Bacillus licheniformis, Bacillus lentus, Bacillus brevis, Bacillus stearothermo-

philus, Bacillus alkalophilus, Bacillus amyloliquefaciens, Bacillus coagulans, Bacillus circulans, Bacillus lautus, Bacillus megaterium, Bacillus thuringiensis, or Streptomyces lividans or Streptomyces murinus, or gram negative bacteria such as E.coli. The transformation of the bacteria may, for instance, be effected by protoplast transformation or by using competent cells in a manner known per se.

PCT/DK99/00088

The yeast organism may favourably be selected from a species of Saccharomyces or Schizosaccharomyces, e.g. Saccharomyces cerevisiae. The filamentous fungus may advantageously belong to a species of Aspergillus, e.g. Aspergillus oryzae or Aspergillus niger. Fungal cells may be transformed by a process involving protoplast formation and transformation of the protoplasts followed by regeneration of the cell wall in a manner known per se. A suitable procedure for transformation of Aspergillus host cells is described in EP 238 023.

In a yet further aspect, the present invention relates to a method of producing a maltogenic alpha-amylase variant of the invention, which method comprises cultivating a host cell as described above under conditions conducive to the production of the variant and recovering the variant from the cells and/or culture medium.

The medium used to cultivate the cells may be any conventional medium suitable for growing the host cell in question and obtaining expression of the maltogenic alpha-amylase variant of the invention. Suitable media are available from commercial suppliers or may be prepared according to published recipes (e.g. as described in catalogues of the American Type Culture Collection).

The maltogenic alpha-amylase variant secreted from the host cells may conveniently be recovered from the culture medium by well-known procedures, including separating the cells from the medium by centrifugation or filtration, and precipitating proteinaceous components of the medium by means of a salt such as ammonium sulfate, followed by the use of chromatographic procedures such as ion exchange chromatography, affinity chromatography, or the like.

### Testing of maltogenic alpha-amylase variants

Maltogenic alpha-amylase variants produced by any of the methods described above may be tested, either prior to or after purification, for amylolytic activity in a screening assay which measures the ability of the variant to degrade starch. The screening in step 10 in the above-mentioned random mutagenesis method of the invention may be conveniently performed by use of a filter assay based on the following procedure: A microorganism capable of expressing the mutated maltogenic alpha-amylase of interest is incubated on a suitable medium and under suitable conditions for secretion of the enzyme, the medium being covered with two filters comprising a protein-binding filter placed under a second filter exhibiting a low protein binding

capability. The microorganism is grown on the second, top filter. Subsequent to the incubation, the bottom protein-binding filter comprising enzymes secreted from the microorganism is separated from the second filter comprising the microorganism. The protein-binding filter is then subjected to screening for the desired enzymatic activity. 5 and the corresponding microbial colonies present on the second filter are identified. The first filter used for binding the enzymatic activity may be any protein-binding filter, e.g., nylon or nitrocellulose. The second filter carrying the colonies of the expression organism may be any filter that has no or low affinity for binding proteins, e.g., cellulose acetate or Durapore™.

Screening consists of treating the first filter to which the secreted protein is bound with a substrate that allows detection of the  $\alpha$ -amylase activity. The enzymatic activity may be detected by a dye, fluorescence, precipitation, pH indicator, IRabsorbance or any other known technique for detection of enzymatic activity. The detecting compound may be immobilized by any immobilizing agent e.g. agarose, agar, 15 gelatine, polyacrylamide, starch, filter paper, cloth; or any combination of immobilizing agents. For example, a-amylase activity can be detected by Cibacron Red labelled amylopectin, which is immobilized in agarose. α-amylase activity on this substrate produces zones on the plate with reduced red color intensity.

10

30

To screen for variants with increased stability, the filter with bound maltogenic 20 alpha-amylase variants can be pretreated prior to the detection step described above to inactivate variants that do not have improved stability relative to the parent maltogenic alpha-amylase. This inactivation step may consist of, but is not limited to, incubation at elevated temperatures in the presence of a buffered solution at any pH from pH 2 to 12, and/or in a buffer containing another compound known or thought to contribute to 25 altered stability e.g., surfactants, EDTA, EGTA, wheat flour components, or any other relevant additives. Filters so treated for a specified time are then rinsed briefly in deionized water and placed on plates for activity detection as described above. The conditions are chosen such that stabilized variants show increased enzymatic activity relative to the parent after incubation on the detection media.

To screen for variants with altered thermostability, filters with bound variants are incubated in buffer at a given pH (e.g., in the range from pH 2-12) at an elevated temperature (e.g., in the range from 50°-110°C) for a time period (e.g., from 1-20 minutes) to inactivate nearly all of the parent maltogenic alpha-amylase, rinsed in water, then placed directly on a detection plate containing immobilized Cibacron Red 35 labelled amylopectin and incubated until activity is detectable. Similarly, pH dependent stability can be screened for by adjusting the pH of the buffer in the above inactivation step such that the parent maltogenic alpha-amylase is inactivated, thereby allowing detection of only those variants with increased stability at the pH in question. To screen

for variants with increased calcium-dependent stability calcium chelators, such as ethylene glycol-bis(ß-aminoethyl ether) N,N,N',N'-tetraacetic acid (EGTA), is added to the inactivation buffer at a concentration such that the parent maltogenic alpha-amylase is inactivated under conditions further defined, such as buffer pH, temperature or a 5 specified length of incubation.

The variants of the invention may be suitably tested by assaying the starchdegrading activity of the variant, for instance by growing host cells transformed with a DNA sequence encoding a variant on a starch-containing agarose plate and identifying starch-degrading host cells as described above. Further testing in regard to altered 10 properties, including specific activity, substrate specificity, cleavage pattern, thermoactivation, thermostability, pH dependent activity or optimum, pH dependent stability, temperature dependent activity or optimum, transglycosylation activity, stability, and any other parameter of interest, may be performed on purified variants in accordance with methods known in the art as described below.

### 15 Degradation of β-limit dextrin by maltogenic alpha-amylase:

Another important parameter in the evaluation of the substrate specificity of maltogenic alpha-amylase variants may be the degree to which such enzymes are capable of degrading starch that has been exhaustively treated with the exoglycosylase B-amylase. To screen for variants which show patterns of degradation on such a 20 substrate differing from the patterns produced by the parent maltogenic alpha-amylase the following assay is performed: β-limit dextrin is prepared by incubating 25 ml 1% amylopectin in McIlvane buffer (48.5 mM citrate and 193 mM sodium phosphate pH 5.0) with 24 μg/ml β-amylase overnight at 30°C. Unhydrolysed amylopectin (i.e., β-limit dextrin) is precipitated with 1 volume 98% ethanol, washed and redissolved in water. 1 25 ml β-limit dextrin is incubated with 18 μl enzymes (at 2.2 mg/ml) and 100 μl 0.2 M citrate-phosphate pH 5.0 for 2 hrs at 30°C and analysed by HPLC as described above. Total hydrolysis of β-limit dextrin is carried out in 2M HCl at 95°C. The concentration of reducing ends is measured by methods known in the art.

### Calcium binding affinity

30

Unfolding of maltogenic alpha-amylases by exposure to heat or to denaturants such as guanidine hydrochloride is accompanied by a decrease in fluorescence, and oss of calcium ions leads to unfolding. Thus, the affinity of a maltogenic alpha-amylase variant for calcium can be measured by fluorescence measurements before and after incubation of the variant (e.g., at a concentration of 10 mg/ml) in a buffer (e.g., 50 mM 35 HEPES, pH 7) with different concentrations of calcium (e.g., in the range from 1 mM-100 mM) or of EGTA (e.g., in the range from 1-1000 mM) for a sufficiently long period of time (such as 22 hours at 55°C).

The measured fluorescence, *F*, is composed of contributions form the folded and unfolded forms of the enzyme. The following equation can be derived to describe the dependence of *F* on calcium concentration ([Ca]):

$$F = [Ca]/(K_{diss} + [Ca])(a_N - b_N \log([Ca])) + K_{diss}/(K_{diss} + [Ca])(a_U - b_U \log([Ca]))$$

where  $a_N$  is the fluorescence of the native (folded) form of the enzyme,  $b_N$  is the linear dependence of  $a_N$  on the logarithm of the calcium concentration (as observed experimentally),  $a_U$  is the fluorescence of the unfolded form and  $b_U$  is the linear dependence of  $a_U$  on the logarithm of the calcium concentration.  $K_{diss}$  is the apparent calcium binding constant for an equilibrium process as follows:

 $K_{diss}$ 

 $N-Ca \ll U + Ca(N = native enzyme; U = unfolded enzyme)$ 

In fact, unfolding proceeds extremely slowly and is irreversible. The rate of unfolding is dependent on calcium concentration, and such dependency for a given enzyme provides a measure of the calcium binding affinity of the enzyme. By defining a standard set of reaction conditions (e.g., 22 hours at 55°C), a meaningful comparison of  $K_{\rm diss}$  for different maltogenic alpha-amylase variants can be made.

20

35

15

### **Industrial Applications**

The maltogenic alpha-amylase variants of the invention possesses valuable properties which may be advantageously used in various industrial applications. In particular, the enzyme finds potential application for retarding or preventing retrogradation, and thus the staling, of starch based food such as common in the baking industry.

The variant may be used for the preparation of bread and other bread products in accordance with conventional techniques known in the art.

It is believed that the modification of the starch fraction by use of the present invention results in increased volume in baked products and improved organoleptic qualities, such as flavour, mouth feel, palatability, aroma and crust colour.

The maltogenic alpha-amylase variant may be used as the only enzyme or as a major enzymatic activity in combination with one or more additional enzymes, such as xylanase, lipase, glucose oxidase and other oxidoreductases, or an amylolytic enzyme.

The enzyme variants of the invention also find industrial applicability as a component in washing, dishwashing and hard-surface cleaning detergent compositions. Some variants are particularly useful in a process for the manufacture of

linear oligosaccharides, or in the production of sweeteners and ethanol from starch, and/or for textile desizing. Conditions for conventional starch conversion processes, including starch liquefaction and/or saccharification processes, are described in, e.g., US 3,912,590 and in EP patent publications Nos. 252,730 and 63,909.

The invention is further illustrated with reference to the following examples which are not intended to be in any way limiting to the scope of the invention as claimed.

## Determination of maltogenic amylase in MANU

One Maltogenic Amylase Novo Unit (MANU) is the amount of enzyme which under standard will cleave one µmol maltotriose per minute. The standard conditions are 10 mg/ml maltotriose, 37°C, pH 5.0, 30 minutes reaction time.

The pH dependence is found by repeating this measurement at the same conditions, but at different pH values.

### **EXAMPLES**

# 15 Example 1: Construction of a variant of Novamyl with altered pH dependent activity

Novamyl is expressed in *Bacillus subtilis* from a plasmid denoted herein as pLBei010. This plasmid contains *amyM* in which the expression of *amyM* is directed by its own promoter and the complete gene encoding Novamyl, e.g., as contained in the strain DSM 11837. The plasmid contains the origin of replication, *ori*, from plasmid pUB110 and an kanamycin resistance marker for selection purposes. pLBei010 is shown in Fig. 1.

### Primer sequences

Site directed mutants of Novamyl were constructed by the megaprimer method essentially as described by Kammann et al. (1989). Briefly, a mutagenic oligonucleotide primer is used together in a PCR reaction with a suitable opposite DNA strand end primer to create a preliminary PCR product. This product is then used as a megaprimer together with another opposite DNA strand end primer to create a double-stranded DNA product. The product of the final PCR reaction was routinely used to replace a corresponding DNA fragment in the pLBei010 plasmid by standard cloning procedures. Mutants were transformed directly into *Bacillus subtilis* strain SHa273, a derivative of *Bacillus subtilis* 168 which is apr, npr, amyE<sup>-</sup>, amyR2<sup>-</sup> and prepared by methods known in the art.

Oligonucleotide primers used in the construction of described variants are as listed below:

Variant Sequence (5'→ 3')

F188H: SEQ ID NO: 3

F188E: SEQ ID NO: 4

F284E: SEQ ID NO: 5

F284D: SEQ ID NO: 6

F284K: SEQ ID NO: 7

N327D: SEQ ID NO: 8

Variant Sequence (3'→ 5') 10

T288K: SEQ ID NO: 9

T288R: SEQ ID NO: 10

Aspartate variants of F284, T288 and N327 were obtained using primer A189 (SEQ ID NO: 11) and B649 (SEQ ID NO: 12) as end-primers.

F188-variants F188L, T189Y were obtained using primer A82 (SEQ ID NO: 13) 15 and B346 (SEQ ID NO: 14) as end-primers.

PCR products with the desired modification(s) were purified, digested with appropriate enzymes, separated by agarose gel electrophoresis and extracted, ethanol precipitated in the presence of glycogen, resuspended in H2O, ligated to pLBei010 20 which had been digested with the same appropriate enzymes, and transformed into Bacillus subtilis SHa273. Transformants were checked for size by colony PCR and for the insertion or removal of specific restriction sites by restriction enzyme digestion. Positive colonies were verified by DNA sequencing methods as described in the art.

### **Fermentation**

25

The B. subtilis SHa273 mutant clones were grown overnight on LB-Kana (10 μg/ml)-Starch plates at 37°C. The colonies from the plate were resuspended in 10 ml Luria broth. One-sixth of each of the suspensions were inoculated into a 500 ml shake flasks containing 100 ml PS-1 media, a soy meal/sucrose-based media, kanamycin for a final concentration of 10  $\mu g/ml$  and 100  $\mu l$  5M NaOH. The pH was adjusted to 7.5 with 30 NaOH before inoculation. The cultures were incubated for five days at 30°C with shaking at 270-300 rpm.

### **Enzyme Purification**

Large particles from the media were removed by flocculation before affinity chromatography. Superfloc C521 (American Cyanmide Company) was used as the 35 cationic flocculant and Superfloc A130 (American Cyanmide Company) as the anionic flocculant.

The culture suspension was diluted 1:1 with deionized water and the pH was adjusted to approx. 7.5. A volume of 0.01 ml of 50<sup>w</sup>/<sub>w</sub>% CaCl<sub>2</sub> per ml diluted culture was added during stirring. A volume of 0.015 ml of 20<sup>w</sup>/<sub>w</sub>% Na-aluminate per ml diluted culture was titrated with 20% formic acid, while keeping the pH between 7 and 8. While stirring 0.025 ml 10<sup>v</sup>/<sub>v</sub>% of C521 per ml diluted culture was added, followed by 0.05 ml 1<sup>w</sup>/<sub>v</sub>% A130 per ml diluted culture, or until flocculation was observed. The solution was centrifuged at 4500 rpm for 30 minutes. Filtration was performed using a filter of pore size of 0.45 µm to exclude larger particles and any remaining bacteria. The filtered solution was stored at -20°C.

### 10 Immobilization of α-cyclodextrin to DSV-agarose

One hundred mg of α-cylcodextrin of molecular weight 972.86g/mol (Fluka 28705) was dissolved in 20 ml coupling buffer (0.5M Na<sub>2</sub>CO<sub>3</sub>, pH 11). Ten ml of DSV-agarose (Mini-Leak, Medium 10-20 mmol/l of divinyl sulfone activated agarose (Kem-En-Tec) was washed thoroughly with deionized water, then dried by suction and transferred to the a-cyclodextrin solution. After the mixture had stirred for 24 hr at ambient temperature, the gel was washed with deionized water, followed by 0.5M KHCO<sub>3</sub>. The gel was transferred to the blocking buffer (20ml 0.5M KHCO<sub>3</sub> + 1ml mercaptoethanol), stirred for 2 hr at ambient temperature, then washed with deionized water.

### 20 Affinity chromatography

The variants were purified by affinity chromatography using the Pharmacia FPLC System. A 0.04 volume of 1M Na-acetate pH 5 was added to the filtrate obtained by flocculation to adjust pH and CaCl₂ was added to a final concentration of 10<sup>-10</sup> M. The solution was filtered and degassed. A Pharmacia XK16 column was prepared with ten ml of the immobilised α-cyclodextrin, then equilibrated in the equilibration buffer (25 mM Na-acetate pH 5) by washing with approximately 10 times the column volume. The filtrate was applied to the XK16 column, which was then washed with the equilibration buffer until protein could no longer be detected in the washing buffer. The column was washed with the equilibration buffer containing 0.5M NaCl to elute nonspecific material, followed by another wash with 2-3 times the column volume of the equilibration buffer. All washings were performed using a flow rate of 10ml/min. Specifically bound material was eluted using a solution of 2% α-cyclodextrin in the wash buffer and collected using the Pharmacia Liquid Chromatography Collector LCC-500 Plus using a flow rate of 5 ml/min.

# Example 2: pH dependent activity of variants

The variants prepared in the preceding Example were tested for activity at various pH values as follows.

A colorimetric glucose oxidase-peroxidase assay for liberated glucose from maltotriose or amylopectin was used to determine the pH activity profiles of the enzyme variants (Glucose/GOD-Perid® Method, Boehringer Mannheim, Indianapolis IN). Activity was assayed in a buffer of 25 mM citrate-phosphate, 0.1mM CaCl<sub>2</sub> at pH values of 2, 2.5, 3, 3.5, 4, 4.5, 5, 5.5, 6, 6.5, 7, 7.5, 8 and 8.6. The buffer pH was adjusted using NaOH and enzymes were diluted in 25 mM citrate-phosphate buffer pH 5. Measurements were taken in duplicate to obtain an average value. All values are relative to the pH at which the highest level of activity is seen.

The results, shown in the table below, indicate that each of the variants has an alteration in the pH dependent activity profile when compared to the parent Novamyl<sup>®</sup>. The highest level of activity for each variant is designated 100% and the activity of that variant measured at the other indicated pH values is a relative percentage of that maximum.

Modifications		рН												
	2.0	2.5	3.0	3.5	4.0	4.5	5.0	5.5	6.0	6.5	7.0	7.5	8.0	8.6
None (parent)	0	0	0	8	47	80	100	95	91	80	66	39	35	30
F188H	1	0	0	1	3	29	77	99	100	88	59	39	31	27
F188E	0	0	0	2	27	62	89	100	93	71	46	28	20	18
T288R	0	0	0	8	51	77	94	100	86	73	50	34	27	12
N327D	1	1	7	27	67	95	100	98	77	33	19	11	5	0

Further, a number of Novamyl variants were tested for activity at pH 4.0 and 5.0, taking the activity of Novamyl at the same pH as 100 %. The activity was determined by hydrolysis of maltotriose (10 mg/ml) at 60°C, 50 mM sodium acetate, 1 mM CaCl<sub>2</sub>. The results are expressed as the ratio between activity at pH 5.0 and pH 4.0:

Modifications	pH 5.0/pH 4.0		
N131D	0.24		
I174Q	0.31		

C207D	0.40
G397P	0.40
H103Y	0.40
∆ <b>262-266</b>	0.47
S32Q	0.53
S32D	0.55
T142A+ D261G	0.62
G370N+ N371G	0.66
S32N	0.68
N176S	0.79
D17E	0.80
None (parent)	1
∆ 191	1.39
192-A-193	1.61
1174E	1.80
192-A-G-193	1.90
∆ <b>192</b>	2.22

The results demonstrate that variants with a higher or lower pH optimum can be obtained according to the invention.

# **Example 3: Thermostability of variants**

# 5 Incubation at 80°C

The thermostability of a number of Novamyl variants was tested by incubating an aqueous solution at 80°C and pH 4.3 and measuring the residual amylase activity at various times. The parent enzyme, Novamyl, was included for comparison. The results are expressed as residual activity at various times in percent of initial activity:

10

Modifications	0	5 min.	10 min.	15 min.	20 min.	25 min.
None (parent)	100	23	9	3	1	0
F188L+ V336L+ T525A	100	63	49	48	52	47
F188I+ Y422F+ I660V	100	71	60	51	43	38
N115D+ F188L	100	73	60	51	44	39

A30D+ K40R+ D261G	100	38	24	15	13	10
T142A+ N327S+ K425E+ K520R+ N595I	100	47	39	25	19	11
F188L+ D261G+ T288P	100	60	67	66	63	67
K40R+ F188L+ D261G+ A483T	100	56	48	40	<b>3</b> 6	30
T288K	100	64	31	18	7	4

The above data show a clearly improved thermostability for the variants compared to the parent amylase.

### Incubation at 85°C with calcium

The Novamyl variant S32E was tested by incubation with 1 mM Ca<sup>++</sup> at 85°C for 15 minutes. The variant showed a residual activity of 48 % whereas the parent enzyme (Novamyl) showed 32 % residual activity at the same conditions.

### DSC

Further, the thermostability was tested for some Novamyl variants by DSC (differential scanning calorimetry) at pH 4.3 or 5.5. Again, the parent amylase was included for comparison. The results are expressed as the denaturation temperature (Tm):

Modifications	Tm at pH 4.3	Tm at pH 5.5
None (parent)	79°C	88°C
N115D+ F188L	86°C	92°C
T142A+ N327S+ K425E+ K520R+ N595I	not determined	93°C

The results show improved thermostability for both variants.

# Example 4: Specific activity of variants

Amylase activity was determined by a colorimetric measurement after action on Phadebas tablets at pH 5.0 and 60°C. The results for two Novamyl variants, relative to Novamyl were as follows:

15

Modifications	Relative amylase activity
None (parent)	100
192-A-193	110
△ (191-195)	300

The specific activity was further tested by action on maltotriose at pH 4.0, 60°C by the MANU method described above. The results showed that the variant G370N, N371G has a maltotriose activity of 106 % compared to Novamyl.

# 5 Example 5: Inhibition of retrogradation

The efficiency of Novamyl and Novamyl variants to inhibit retrogradation was determined as follows:

730 mg of 50 % (w/w) amylopectin slurry in 0.1 M sodium acetate, at a selected pH (3.7, 4.3 or 5.5) was mixed with 20 µl of an enzyme sample, and the mixture was incubated in a sealed ampoule for 1 hour at 40°C, followed by incubation at 100 °C for 1 hour in order to gelatinize the samples. The sample was then aged for 7 days at room temperature to allow recrystallization of the amylopectin. A control without enzyme was included.

After aging, DSC was performed on the sample by scanning from 5°C to 95°C at a constant scan rate of 90°C/hour. The area under the first endothermic peak in the thermogram was taken to represent the amount of retrograded amylopectin, and the relative inhibition of retrogradation was taken as the area reduction (in %) relative to the control without enzyme.

In the table below, the efficiency of the enzyme is expressed as the ratio of the relative inhibition of retrogradation to the enzyme dosage (in MANU/ml):

рН	Modifications	MANU/ml	Relative inhibition	Efficiency
3.7	A30D+ K40R+ D261G	0.23	0.38	1.7
3.7	T142A+ N327S+ K425E+ K520R+ N595I	0.07	0.29	4.1
3.7	None (parent)	0.27	0.38	1.4
4.3	N115D+ F188L	0.01	0.18	18
4.3	None (parent)	0.27	0.43	1.6

5

10

5.5	△ (191-195)+ F188L+ T189Y	0.02	0.12	6
5.5	∆ (1 <b>91-195</b> )	0.02	0.14	7
5.5	∆ <b>(191-195)</b>	0.05	0.31	6.2
5.5	N115D+ F188L	0.01	0.39	39
5.5	T142A+ D261G	0.14	0.53	3.8
5.5	None (parent)	0.27	0.49	1.8

The results demonstrate that a number of variants are more efficient than the parent amylase to inhibit retrogradation.

# Example 6: Anti-staling effect of variants

Bread was made by the European Straight Dough method or from sour dough with or without addition of enzymes, and loaves were baked in lidded pans, to avoid volume effects. The bread was allowed to cool for 2 hours, and the texture was analyzed. The remaining loaves were then wrapped in plastic bags and stored at room temperature for texture analysis after 1, 4 and 7 days.

The texture analysis of each loaf was done by cutting 4 slices; the force was measured at 25 % compression (P1), at 40% compression (P2) and after keeping 40% compression constant for 30 sec. (P3). P1 was taken as the firmness, and the ratio (P3/P2) was taken as the elasticity of the crumb. The extent of retrogradation after 7 days storage was determined by DSC as described in Example 7.

### 15 European Straight Dough (pH5.5-6.0)

A Novamyl variant (T142A+ N327S+ K425E+ K520R+ N595I) was tested at dosages in the range of 0-2 mg enzyme/kg flour, and the parent enzyme (Novamyl) was used for comparison.

The results showed that at equal dosages, the variant gives better elasticity (P3/P2) than the parent enzyme after two hours and 1 day. The results after 7 days showed that the variant at dosages of 1-2 mg/kg gave significantly softer crumb (lower firmness, P1) than the parent enzyme at the same dosage. Thus, the variant has a better anti-staling effect throughout a 7-day storage period.

### Sour dough (pH approx. 4.5)

A Novamyl variant (F188L+ D261G+ T288P) was tested in sour dough, and the parent enzyme (Novamyl) was used for comparison. The following results were

obtained for firmness (P1) after 7 days, elasticity (P3/P2) after 4 and 7 days and ret-

rogradation after 7 days:

Enzyme	Dosage mg/kg flour	Firmness (P1) after 7 days
None	0	2590
	1	2031
Parent	3	1912
	13	1570
Variant	1	1436
	3	1226

Enzyme	Dosage mg/kg flour	Elasticity 4 days	Elasticity 7 days
None	0	0.49	0.47
	1	0.51	0.52
Parent	3	0.53	0.51
	13	0.53	0.51
Variant	1	0.59	0.57
	3	0.57	0.58

Enzyme	Dosage mg/kg flour	Retrogradation, 7 days (relative to control)
None	0	100 %
	1	100 %
Parent	3	63 %
	13	32 %
Variant	1	46 %
	3	20 %

The results show that the variant has a markedly improved effect on texture evaluated as firmness and elasticity in sour dough at pH 4.5. A dosage of 1-3 mg/kg of the variant is superior to 13 mg/kg of the parent enzyme on all parameters tested, and the elasticity achieved with the variant cannot be matched by the parent enzyme at any dosage.

## Example 7: Cleavage pattern of variants

The cleavage pattern in starch hydrolysis was compared for two variants and the parent enzyme, Novamyl.

The results below indicate % by weight of each oligosacccharide (G1-G8) formed after 24 hours incubation in 1% (w/v) starch using 50 mM sodium acetate, 1 mM CaCl<sub>2</sub>, pH 5.0 at 50 °C. The oligosaccharides were identified and quantitated using HPLC.

Oligosaccharide	Parent	∆ (191-195)	N115D+ F188L
G8	-	1.7	-
<b>G7</b>	-	2.6	-
G6	-	7.5	1.4
<b>G</b> 5	-	10.1	2.1
G4	-	21.1	11.3
<b>G</b> 3	-	28.7	10.7
G2	96.5	28.3	61.9
G1	3.5		12.6

The results demonstrate a significantly altered cleavage pattern. Novamyl after 24 hours produces mainly maltose and virtually no higher oligosaccharides. In contrast, the two variants produce significant amounts of maltotriose and higher oligosaccharides.

#### **REFERENCES CITED**

Klein, C., et al., *Biochemistry* 1992, **31**, 8740-8746,

Mizuno, H., et al., *J. Mol. Biol.* (1993) **234**, 1282-1283,

Chang, C., et al, *J. Mol. Biol.* (1993) **229**, 235-238,

Larson, S.B., *J. Mol. Biol.* (1994) **235**, 1560-1584,

Lawson, C.L., *J. Mol. Biol.* (1994) **236**, 590-600,

Qian, M., et al., *J. Mol. Biol.* (1993) **231**, 785-799,

15

25

Brady, R.L., et al., Acta Crystallogr. sect. B, 47, 527-535,

Swift, H.J., et al., Acta Crystallogr. sect. B, 47, 535-544

A. Kadziola, Ph.D. Thesis: "An alpha-amylase from Barley and its Complex with a Substrate Analogue Inhibitor Studied by X-ray Crystallography", Department of Chemistry University of Copenhagen 1993

MacGregor, E.A., Food Hydrocolloids, 1987, Vol.1, No. 5-6, p.

B. Diderichsen and L. Christiansen, Cloning of a maltogenic  $\alpha$ -amylase from Bacillus stearothermophilus, FEMS Microbiol. letters: 56: pp. 53-60 (1988)

Hudson et al., Practical Immunology, Third edition (1989), Blackwell Scientific Publications,

Sambrook et al., <u>Molecular Cloning: A Laboratory Manual</u>, 2nd Ed., Cold Spring Harbor, 1989

S.L. Beaucage and M.H. Caruthers, <u>Tetrahedron Letters</u> <u>22</u>, 1981, pp. 1859-1869

Matthes et al., The EMBO J. 3, 1984, pp. 801-805.

R.K. Saiki et al., Science 239, 1988, pp. 487-491.

Morinaga et al., (1984, Biotechnology 2:646-639)

Nelson and Long, Analytical Biochemistry 180, 1989, pp. 147-151

Hunkapiller et al., 1984, Nature 310:105-111

20 R. Higuchi, B. Krummel, and R.K. Saiki (1988). A general method of *in vitro* preparation and specific mutagenesis of DNA fragments: study of protein and DNA interactions. *Nucl. Acids Res.* **16**:7351-7367.

Dubnau et al., 1971, J. Mol. Biol. 56, pp. 209-221.

Gryczan et al., 1978, J. Bacteriol. 134, pp. 318-329.

S.D. Erlich, 1977, Proc. Natl. Acad. Sci. 74, pp. 1680-1682.

Boel et al., 1990, Biochemistry 29, pp. 6244-6249.

Kammann, M Laufs, J Schell, J and Gronnenborn, B (1989) Nucleic Acids Research 20:4937-4938.

9.80 11.138 11.138 11.138 11.130 11.1 26.131 26.904 28.574 26.561 25.099 25.099 25.184 24.863 25.709 23.953 24.863 25.709 23.953 24.863 25.709 23.953 24.863 20.852 21.970 25.646 46.350 44.425 44.425 45.577 46.166 45.340 44.293 44.293 43.319 43.442 28.343 24.346 23.831 21.755 26.527 22.675 27.608 64.488 64.488 56.488 56.982 55.015 55.015 56.096 56.097 60.014 60 578 18.160 17.698 21.036 22.11.036 23.133 24.040 37.106 40.113 39.826 40.643 40.537 40.931 42.135 **aaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaaa** LLYS
GGLYY
GGLY
GGLYY
GGLY
GGLYY
GGL 16.73 26.53 21.07 12.71 13.38 112.57 112.57 116.30 116.30 116.30 116.30 116.30 116.30 116.30 116.30 117.81 10.01 36.680 37.316 37.029 35.219 39.355 38.956 40.380 38.175 37.898 38.726 39.369 36.045 36.045 38.551 39.209 39.209 38.123 34.568 38.384 41.357 43.843 41.427 12.785 56.595
55.462
56.773
56.773
56.773
56.773
56.773
56.773
57.682
57.682
57.682
57.682
57.682
57.682
57.682
57.682
57.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682
67.682 656 349 604 360 560 588 10.254 11.216 12.585 11.527 11.527 13.05 13.05 14.06 15.09 15.09 15.09 15.09 15.09 15.09 15.09 15.09 16.09 1 19.340 20.589 21.169 22.349 21.639 21.002 21.874 20.369 20.369 21.508 z 5 u o 8 8 z 5 u o 8 8 z 5 u o 8 8 z 5 u o 8 8 z 5 u o 8 8 z 5 u o 8 8 2 5 u o 8 8 2 5 u o 8 8 8 

45

15.02 116.37 116.40 116.40 117.99 116.04 117.99 117.99 113.05 113.05 117.99 117.10 117.10 117.30 117.46 117.30 117.30 117.30 13.475 14.010 13.739 14.802 14.471 14.772 14.825 15.246 16.739 17.224 17.497 14.890 15.290 14.189 14.499 15.662 17.001 12.920 11.854 18.083 17.340 15.923 15.252 17.367 16.673 18.772 15.423 14.019 13.724 12.538 46.150 47.743 47.743 48.366 45.532 43.924 44.924 49.871 50.862 50.103 49.981 50.066 49.796 50.401 51.353 52.337 53.461 50.575 49.884 49.932 51.954 52.918 52.705 52.148 52.497 53.538 54.685 53.086 37.681 38.769 38.769 39.124 40.033 37.228 37.228 36.349 36.215 36.215 36.248 36.215 36.248 36.248 36.248 36.248 37.248 37.248 37.248 37.248 37.248 37.248 37.35 37.36 37 39.211 39.604 38.672 39.059 41.036 44444444444444444444444444444444444444 14.05 15.80 16.74 19.32 19.70 22.69 13.46 14.61 13.84 12.67 15.87 27.943 27.800 25.321 27.719 27.504 27.214 27.062 24.586 23.134 .097 22.413 425 23.040 23.110 215 261 49.476 49.813 48.696 47.270 46.968 46.285 44.982 43.422 48.893 47.226 47.226 48.665 48.444 45.934 45.410 44.264 53.057 51.614 51.024 53.294 53.294 52.543 52.947 52.187 50.019 51.054 69.440 69.476 47.219 288 450 48. 296 543 798 798 798 798 790 184 154 154 730 37.927 39.216 37.160 37.160 39.717 37.658 38.935 38.662 38.414 37.754 38.426 39.678 39.678 565 759 430 683 683 786 505 570 570 4444444444444444444444444444444 

46 14.18 110.53 112.89 113.47 113.43 113.00 115.18 116.88 117.90 117.90 117.90 117.90 117.90 118.65 118.65 119.66 119.66 119.60 119 17.122 16.416 17.135 17.441 17.463 17.991 19.028 16.821 17.548 19.436 16.450 58.755 59.737 60.460 61.033 61.613 61.613 61.613 61.613 62.003 63.492 63.492 63.492 63.492 63.492 63.492 63.492 63.492 63.492 63.492 63.492 63.493 63 51.843 53.192 53.031 53.080 52.024 52.056 51.189 50.946 53.773 53.847 42.472 40.784 40.170 38.783 41.090 43.025 44.333 42.749 42.749 43.317 43.317 44.382 44.382 43.317 44.382 44.382 44.382 44.382 46.646 46.700 46.700 46.700 48.507 48.543 49.278 49.278 49.278 49.278 49.375 48.513 48.367 47.570 49.385 50.469 44444444444444444444444444444444444444 15.54 16.99 19.06 19.06 19.27 18.27 115.11 14.64 14.25 113.99 113.23 114.25 115.05 117.62 117 11.167 8.010 8.045 8.444 7.818 8.050 8.364 9.038 10.193 10.460 9.161 53.483 53.615 53.841 53.244 54.024 55.375 56.355 53.268 55.514 56.812 56.812 59.094 56.667 57.569 58.766 57.610 58.587 57.989 57.988 57.968 58.851 57.664 57.524 56.801 57.676 58.848 56.295 55.254 55.618 53.027 54.670 53.409 52.483 57.105 57.836 58.672 35.087 36.3097 36.3097 36.3097 36.3097 36.3097 36.3097 37.3098 37.3097 41.166 44.235 42.408 43.281 \* 

47

15.83 11.5.83 11.3.24 11.3.26 11.3.26 11.3.26 11.3.26 11.5.30 11.5.30 11.50 22.334 22.068 19.114 18.112 19.768 19.768 19.495 10.692 11.693 11 17.660 17.416 16.436 18.105 16.146 20.188 579 881 547 711 077 086 037 820 603 715 484 68.171
69.681
60.686
61.576
60.686
60.686
60.686
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60.710
60 45.254 44.44 44.44 44.125 45.125 45.125 45.125 45.125 45.125 45.125 45.125 45.125 46.125 17.78 19.03 18.65 22.34 22.34 22.34 16.85 16.85 17.21 16.94 17.21 18.69 19.77 112.77 113.20 113.20 114.00 11.08 11 23.685 23.207 24.031 23.507 23.507 23.590 21.108 25.364 26.297 26.617 27.501 27.814 28.265 56.348 54.324 52.54324 52.54333.325 52.5433.325 52.5433 52.5433 52.5433 52.5433 52.5433 52.5433 52.5433 52.5433 52.5433 60.0250 60.0 50.009
44.104
44.1023
45.104
46.535
46.1033
47.1033
47.1033
47.1033
47.1033
47.1033
47.1033
47.1033
47.1033
47.1033
47.1033
47.1033
47.1033
47.1033
47.1033 CLYSSER SERVES S 

48

14 84 84 13 13 64 64 67 67 67 28.101 28.652 29.596 26.819 27.165 25.358 25.358 25.358 26.233 30.611 27.536 29.945 30.611 27.536 27.536 28.23 30.611 27.536 28.23 30.611 27.536 30.611 37.536 37.5 31.625 30.818 31.318 30.729 33.338 34.668 35.732 32.723 31.261 32.388 30.943 34.670 35.937 44.040
42.905
42.905
42.905
44.904
44.317
44.3139
43.318
43.318
43.318
43.318
43.318
44.339
43.318
44.339
43.318
45.574
46.651
46.752
47.112
46.752
47.112
46.753
47.112
47.112
48.339
48.3360
48.33960
48.33960
48.33960
48.33960
48.33960 46.366 48.136 44.107 43.396 44.139 46.878 29.382 30.1112 29.684 28.627 27.364 27.364 27.364 27.364 27.364 27.365 29.125 33.265 3 12.42 12.08 9.81 10.92 10.92 10.92 11.80 11.80 11.19 11.19 11.19 11.10 1 16.49 12.03 13.18 12.67 11.91 20.845 22.221 23.230 22.614 21.880 23.065 23.065 22.429 23.344 24.489 21.094 21.327 22.384 20.532 22.778 23.617 23.973 25.090 22.869 22.638 21.666 23.954 23.037 23.443 25.382 25.382 22.122 22.22.295 22.992 23.005 23.419 24.628 25.680 27.005 28.000 .005 28.280 28.715 29.879 28.220 51.215 52.862 52.862 52.862 53.629 51.085 51.085 51.085 51.086 51.086 51.086 51.086 51.086 64.090 48.111 50.947 51.255 52.143 47.952 38.208 38.834 37.789 36.451 36.173 37.178 37.178 37.136 37.136 37.148 34.69 37.148 30.227 30.680 33.114 34.108 33.114 33.737 33.737 31.096 31.055 31.055 31.055 31.055 31.055 31.055 

																					4	9																			
ω	7	7	9	9	ω	9	9	9	9	7	9	9	89	7	9	9	æ	9	9	9	7	9	9	ω.	9	æ	9	7	9	9	89	9	œ	9	7	9	9	æ	9	9	9
0.33 14.24				1.00 14.21	1.00 14.81	1.00 14.15	1.00 13.04	1.00 12.46	1.00 12.70	1.00 14.23	1.00 13.78	1.00 14.56	1.00 13.13					1.00 11.10	1.00 12.12		1.00 10.70		1.00 13.77						_		11.8	13.8	73	1.00 14.32		1.00 8.17					1.00 11.22
41.279	43.537	41.498	42.087	42.837	43.659	40.930	41.274	42.303	39.963	42.566	43.205	42.311	42.758	41.014	40.114	39.842	39.617	38.744	37.731	38.974	39.992	39.675	38.565	38.103	40.898	41.874	41.584	38.156	36.914	36.193	36.775	37.236	37.793	35.965	34.896	34.073	33.377	32.734	32.986	33.736	31.900
42.416	42.519	47.646	48.909	49.656	50.516	49.777	51.201	51.132	51.825	49.492	50.247	51.337	52.22	51.265	52.267	52.007	50.836	52.192	53.220	52.489	53.016	52.822	53.737	53.538	52.926	53.791			'n		56.214	56.837	56.649	57.734	55.287	55.600	56.897	57.080	54.510	53.214	54.893
34.480	34.884	32.481	32.993	31.893	32.253	33.536	34.050	35.177	34.587	30.605	29.537	28.987	28.207	29.343	28.773	27.297	26.933	29.491	28.892	30.961	26.431	25.022	24.526	23.404	24.072	24.680	24.085	25.351	25.042	26.379	27.313	24.388	23.094	24.269	26.394	27.605	27.249	26.200	27.714	28.160	28.737
63	63	64	64	64	64	64	64	64	64	65	65	9	65	99	99	99	99	99	99	99	67	29	29	67	67	29	29	68	89	68	68	68	68	89	69	69	69	69	69	69	69
OE1BGLN A		LEU A		LEU A	LEU A	B LEU A			CD2 LEU A	GLY A	CA GLY A	C GLY A	GLY A			VAL A						A THR A		THR A		THR	~		CA THR A			B THR A		~	N ILE A	A ILE A	ILE	ILE			CG2 ILE A
_	-		1 1		3 0	4 CB				8																															_
20	509	510	51	51	513	514	515	516	517	51	519	520	521	522	52	52	52	52	52	52	52	530	53	53	533	53	53	53	537	53	539	540	541	542	543	544	545	546	547	548	54
9	7	9	9	80	9	9	9	9	9	9	9	œ	7	9	9	80	9	9	9	9	7	9	9	80	9	9	9	9	7	7	9	9	æ	9	9	9	œ	7	9	9	9
1.00 16.71		1.00 12.47			1.00 12.16	1.00 12.17	1.00 13.23	1.00 12.24	1.00 12.99	1.00 14.93	1.00 14.75	1.00 13.93	1.00 11.74	1.00 11.91	1.00 14.11	1.00 13.33	1.00 12.06	1.00 13.18	1.00 15.35	1.00 16.34	1.00 12.47		1.00 13.60	1.00 15.97	1.00 14.71	1.00 17.27	1.00 30.17	1.00 25.52	1.00 24.63	1.00 12.05		1.00 13.71	1.00 13.66	0.66 17.93	0.66 27.01	0.66 35.95	.66 4	0.66 36.04	0.33 7.59	8.2	•
33,514	36.210	36.994	37.584	38.797	36.026	36.667	37.462	36.453	38.049	37.022	37.823	38.464	36.735	37.225	38.272	39.283	36.051	34.992	33.753	35.605	38.162	39.202	40.535	41.576	38.730	39.764	39.223	40.343	40.515	40.455	41.660	42.325	43.544	41.262	40.865	40.343	41.149	39.043	41.345	42.621	42.408
43.247		45.511		46.994	45.920	46.938	46.528	48.293	47.509	49.268	48.852	49.782	47.410	48.610	48.316	49.022	49.382	49.809	50.310	50.971	47.217	46.844	46.558	47.032	45.588	45.263	44.363	44.088	45.203	45.781	45.448	46.705	46.751	44.501	43.092	42.362	42.022	42.128	44.387	43.793	42.824
30,807	32.806	33.789	33.072	33.237	34.918	35.856	36.917	35.602	37.730	36.438	37.473	38.287	32.298	31.622	30.570	30.508	30.993	32.030	31.263	32.865	29.850	28.890	29.614	29.149	28.117	27.011	25.908	24.879	23.887	30.722	31.437	32.010	32.200	32.582	32.178	33.421	34.283	33.525	32.511	33.072	4.23
9	9	9	9	9	09	09	9	9	9	09	09	9	61	61	61	61	61	61	61	61	62	62	62	62	62	62	62	62	62	63	63	63	63	63	63	63	63	63	63	63	63
A CAI	TYR	CA TYR A	TYR	O TYR A	3 TYR	TYR	1 TYR	CD2 TYR A	TYR	TYR	TYR		LEU		LEU	LEU	LEU		LEU	LEU		LYS	LYS		3 LYS		LYS	CE LYS A	LYS	N GLN A	CA GLN A	C GLN A	O GLN A	CB AGLIN A	AGLN					CG BGLN A	BGLN
17.1	472	473	474	475	921	477	478	479	480	481	482	483	484	485	486	487	488	489	490	491	492	493	494	495	496	497	498	499	200	501	502	503	504	505	905	507	508	509	505	206	507

8.87 12.04 9.81 9.30 11.34 13.04 9.32 12.70 14.30 14.19 15.14 12.50 12.26 15.155 13.979 12.745 12.895 11.641 13.385 16.619 17.945 18.600 12.866 13.026 15.534 62.605 63.683 61.153 61.079 60.856 64.886 65.195 63.977 63.257 66.197 58.894 60.912 61.136 61.464 61.678 62.279 61.966 63.178 60.864 61.296 61.292 62.605 .239 67.597 67.439 63.867 570 759 370 988 636 62.042 947 691 670 63. 61. 62. 63. 64. 63 25.882 27.708 27.108 28.305 29.436 26.111 24.952 24.074 27.958 29.020 29.821 29.020 29.821 29.171 31.131 31.131 31.131 32.764 33.766 34.766 34.766 34.766 35.766 36.766 36.766 36.766 36.766 37 35.393 36.754 37.275 34.801 34.468 13.98 10.83 10.49 11.31 10.98 10.17 10.33 10.62 8888 28.240 27.096 28.066 27.218 31.118 30.225 28.276 27.923 28.322 572 26.237 .048 .042 28.439 944 25.920 25.128 25.358 63.016
63.016
64.729
64.729
64.729
64.234
65.061
58.311
58.311
59.311
56.159
56.159
56.169
60.998
60.011
60.898
60.018
60.018
60.018
60.018
60.018
60.018 313 286 538 338 27.937 28.196 28.095 28.991 30.214 28.954 29.050 29.645 30.017 29.645 30.017 29.645 30.017 29.645 30.017 29.645 30.017 29.645 30.017 29.645 30.017 29.645 30.017 30 447 427 521 653 672 345 351 998 230. **ARARARARARARARARARARA** TILE
TIRP P
TIRP 

51 111.19 110.24 110.27 110.21 110.03 110.03 111.70 111.70 111.07 111.07 111.90 110.92 110.92 110.92 110.92 110.92 110.92 110.92 110.92 110.92 110.92 110.92 110.92 110.92 110.92 110.93 110.92 110.93 11 9.73 8.93 10.94 10.18 9.66 9.79 9.79 11.09 00.1 15.375 21.732 21.106 21.748 17.677 18.341 18.063 16.705 16.251 17.317 18.471 18.124 17.887 18.721 19.121 19.087 955 21.412 22.115 23.777 23.443 24.411 507 926 906 010 995 837 555 .404 62.467 64.357 64.329 63.063 60.489 60.769 61.653 62.857 61.256 63.68 62.054 64.126 64.126 62.703 62.983 63.104 68.295 68.295 67.535 69.546 65.099 64.459 64.643 65.021 62.712 62.404 61.241 61.476 64.338 65.403 66.903 66.862 035 898 44.190
44.140
44.140
44.141
44.141
44.141
44.141
44.141
44.141
44.141
44.141
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44.161
44 112.91 11.11 14.40 12.89 11.52 12.02 11.47 11.91 11.91 11.33 13.989 10.830 9.927 12.244 12.629 12.798 13.202 14.316 12.484 12.923 13.060 13.230 13.444 15.687 12.873 14.019 17.030 15.747 15.275 14.496 13.888 14.074 14.933 64.479 64.930 66.930 66.937 66.927 66.927 67.287 67.888 64.345 64.345 64.345 67.888 67.213 69.432 68.477 67.298 65.652 64.552 64.902 64.416 64.077 63.600 63.065 63.782 65.712 38.652 39.521 39.072 38.583 38.583 38.265 39.072 39.072 44.059 44.051 44.053 44.053 44.053 44.053 44.053 44.053 44.053 44.053 44.053 44.053 44.053 44.053 44.053 44.054 45.054 46 50.533 47.462 47.117 45.986 48.356 49.500 49.284 50.645 45.024 44.024 42.644 

534 CA THR
535 C THR
536 O THR
639 CG THR
640 N LEU
641 CA LEU
641 CA LEU
642 CA LEU
644 CB LEU
644 CB LEU
645 CG LEU
646 CD LEU
647 CD LEU
647 CD LEU
648 N ALA
649 CA ALA
655 CA ALA
655 CA ALA
655 CA ALA
655 CA ALA
656 CA ALA
656 CA ALA
657 N THR
658 CA THR
658 CA THR
659 CA THR
659 CA THR
659 CA THR
659 CA THR
650 CA THR
651 CB THR
652 CB THR
653 CB THR
653 CB THR
654 N ASP
655 CA ASP
655 CA ASP
656 CA ASP
657 N ASP
657 N ASP
658 CA ASP
658 CA ASP
659 CA ASP
659 CA ASP
659 CA ASP
650 CA ASP
651 CA ASP
651 CA ASP
652 CA ASP
653 CA ASP
653 CA ASP
654 N ASP
655 CA ASP
655 CA ASP
656 CA ASP
656 CA ASP
657 CA ASP
657 CA ASP
657 CA ASP
658 CA ASP
658 CA ASP
659 CA ASP
650 CA ASP

11. 15. 12 11. 44 11. 144 11. 10 12. 67 14. 77 11. 10 15. 82 16. 91 17. 10 18. 29 19. 98 1 17.373 16.982 17.422 16.532 18.799 17.067 19.296 15.152 14.982 14.991 13.835 13.915 15.036 14.853 15.625 11.187 63.426 63.125 65.844 66.935 67.807 68.002 68.073 68.93 68.333 61.786 60.424 59.424 61.999 63.337 61.999 64.005 64.005 58.975 58.962 57.830 64.026 65.176 64.640 64.166 63.154 63.467 63.012 63.251 .615 639 330 581 26.880 25.536 24.977 25.448 23.229 24.238 24.238 24.238 24.238 24.238 24.238 24.238 24.238 24.238 25.209 27.248 15.96 10.95 9.91 10.45 9.91 11.32 9.84 19.345 17.415 19.258 18.020 18.932 19.040 18.171 16.661 17.495 17.079 12.945 21.753 20.459 20.520 22.837 24.219 24.730 26.247 26.247 26.715 17.103 19.694 15.074 18.891 14.292 67.152 67.935 69.141 67.935 67.385 67.713 67.147 65.888 66.237 68.040 67.343 68.040 67.343 68.057 68.059 68.059 68.059 68.059 69 32.653 32.556 33.132 33.132 33.217 34.112 31.905 31.905 32.829 32.829 32.914 31.949 32.914 32.914 31.949 32.914 31.949 32.918 32.918 33.956 36.448 37.359 36.448 37.359 37 28.643 27.263 29.315 28.819 27.382 26.834 29.766 29.739 30.894 

21.053 17.759 17.759 11.566 11.5927 11.5927 11.5928 11.749 11.749 11.992 11.949 11.969 58.005
52.584
551.533
552.143
552.143
552.143
552.143
552.144
552.156
652.169
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104
673.104 34.365 30.491 30 11. 20 13. 05 13. 05 11. 0. 14 11. 0 19. 385 19. 486 19. 486 10. 55.834 56.844 57.844 26.080
27.229
28.6029
28.6029
29.6029
29.6029
30.918
30.918
30.918
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192
31.192 1000 11000 11000 11000 11001 11001 11001 11002 11003 1003 

12.99 15.11 15.19 12.99 12.99 12.99 12.94 12.94 12.94 13.19 13.19 13.19 13.19 13.19 14.86 11.99 28.961 29.987 24.868 24.868 22.6633 22.6633 23.936 31.933 31.933 32.766 33.753 33.753 33.131 33.131 33.131 33.131 33.131 33.131 33.133 33.763 34.763 36.763 36.763 36.763 36.763 36.763 36.763 36.763 36.763 36.763 36.763 36.763 935 620 781 694 46.553 46.160 47.261 47.261 47.261 47.261 47.261 48.289 48.289 48.289 48.289 48.289 48.289 48.399 48.208 48.399 59.399 48.399 59 44.412 43.445 20.678
20.121
19.372
118.200
118.200
118.200
119.061
117.304
22.731
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
23.083
20.028
20.028
20.028
20.028
20.028
20.028
20.028
20.028
20.028
20.028
20.028
20.028
20.028
20.028
20.028
20.0380
20.0380
20.0380
20.0380
20.0380
20.0380
20.0380
20.0380
20.0380
20.0380
20.0380
20.0380
20.0380
20.0380
20.0380
20.0380 **AAAAAAAAA** 13.20 13.20 12.63 14.41 14.41 13.70 13.38 14.48 14.48 15.04 15.04 15.04 15.04 11.25 11.25 11.25 11.35 11.35 11.35 11.35 11.35 11.35 11.35 11.35 11.35 11.35 11.35 11.35 11.35 11.35 11.35 11.35 22 . 126 22 . 830 24 . 060 21 . 525 22 . 611 23 . 252 22 . 913 24 . 247 23 . 876 24 . 558 22 . 065 22 . 065 22 . 755 23 . 755 27.037 28.040 26.664 26.823 27.762 28.384 24.698 21.610 20.937 21.305 21.305 22.997 23.718 25.017 25 25.811 28.403 .512 51.800
51.137
51.137
51.137
52.1036
54.128
54.128
55.850
55.850
55.858
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
64.993
65.993 24. 247 23. 058 23. 30 061 23. 058 23. 30 061 23. 30 061 23. 30 061 24. 30 061 24. 30 061 25. 30 06 \$\\ \text{2} \\ \t 

WO 99/43794	PCT/DK99/00088
110 22/10/21	

voα	<b>ω</b> (	۰		۰ ۲	9	9	<b></b>	ω (	ے م	. 00	7	9	9	80	9 1	<b>9</b> 4	o vo	9	55	9 1	۰ ۰	9	<b>.</b>	φ,	ی م	7	9	9	œ	9 '	י מ	٦ ۵	۰ ب	9	89
1.00 12.61		1.00 11.44	7 [					1.00 9.91	1.00 11.94			00.	8			1.00 8.93		1.00 12.04		1.00 10.20	1.00 9.78				1.00 10.62		00	11.3		2 :	12	1 00 0 48	10	11.6	
	28.254	26.823	26.856	27.77					28.135			25.077	24.887			24.223				22.070	23.398				23.590	855	21.986				24.189	24.203			881
62.207	63.080	60.356	58.883	62.445	63.870	64.374	63.710	63.719	64.951 66.041	64.880	65.484	65.952	67.394	68.163	900.99	66.405	67.627	66.014	67.993	67.218	68.932	69.845	70.026	68.381	67.246	70.638	71.474	72.802	73.565	71.673	71.602	*TC.0/	74.481	74.637	75.724
25.247	24.413	24.039	23.704	26.535	7.	26.509	26.837	5 52	29.305	30.335	25.802	25.134	25.609	25.752	23.609	22.760	22.095	21.907	21.325	21.229	26.851	26.171	26.494	28.178	28.829	25.165	24.341	an i	24.336	22.983	23.480	0 -	26.687	27.244	27.256
VAL A 126 VAI. A 126	⋖ •	VAL A 126	( A	: ∢	ø	Æ	A.	ASP A 127		: A	Ø	PHE A 128	Ø	Ø	PHE A 128		£ 4	PHE A 128	Æ	PHE A 128	£ 4		Æ.		VAL A 129 VAL A 129	Ø	K	Ø	Ø			£ 4	: 4	Ø	Ø
υο	0 8	9 8	3 8	2	ජ	ပ	0	8	3 5	007	z	ర	U	0	පි දි	9 8	3 8	8 8	CE2	Z Z	ූ ජි		0	<u>m</u> i	3 8	z	చ	ပ	0	ප (	9 8	3 2	: ව්	ບ	0
1012	1013	1014	1016	1017	1018	1019	1020	1021	1022	1024	1025	1026	1027	1028	1029	1030	1032	1033	1034	1035	1037	1038	1039	1040	1041	1043	1044	1045	1046	1047	1048	1040 1040	1051	1052	1053
vov	9 (	æι		. 6	9	œ	7	۰ ب	φα	o vo	9	9	9	7	9	ه م	ט מ	w	9	9 1	, ,	· w	9	œ <b>ч</b>	ہ م	v								7	
	00 20	1.00 23.34	73.6	16.9	13	00		8	1.00 13.96		8		00		1.00 11.74		1.00 11.17	1.00 12.23	1.00 16.30	1.00 21.38	1.00 9.54	1.00 9.53	1.00 12.31	1.00 12.23	1.00 12.71	1.00		1.00	1.00		1.00	1.00	1.00	1.00	-
35.481	34.799	33.576	120.05	· .	37.772	38.747	36.527	36.322	35.289	34.22	37.040	35.506	36.465	35.609	34.652	33.786	35.457	36.163	36.815	38.025	32.656	31.740	31.460	30.990				31.352			32.47	33.069	1 6	0,	7
ν	<u>ښ</u> ،	43.805	46.312		49.129	49.890	49.397	50.569	51.454	50 129	49.424	51.348	48.709	52.752	53.718	54.362	54 811	54.205	55.310	56.014	54.936	55.533	56.954	57.168	41	53.313						156.00	3 5	9	9
24.337	24.753	24.778	9/0.52	20.994	21.840	21.866	22.262	23.128	22.464	24 556	25.320	25.415	26.569	22.344	21.767	22.865	24.052	19.832	18.994	19.601	22.372	23.343	22.856	21.723	23.372	23.875	23.726	23.419	24.430	25.549	23.403	24.811	24.770	23.971	24.864
ASN A 120	Æ.	<b>A</b>	ASN A 120	4	Ø	æ	æ	4	ILE A 122	ζ 4	< <	A	ILE A	æ	Æ,	< ⋅		LYS A 123	æ		LIS A 123 VAT. A 124	VAL A 124	VAL A 124	VAL A 124	'B VAL A 124	VAL A	ILE A	ILE A	æ	ILE A	ILE A	CG1 1LE A 125	4 4 4	VALA	VAL A
88			NDZ	z 5					o c	ء د	-	CG2	~		ð		ع و			8		: 5	U		_ (	9 0	z	S	U	_	8	9 (	ככ	) z	: 5

56 9 13.44 115.17 110.70 110.33 110.13 110.33 110 44 28 28 38 38 08 63 13.738 11.779 13.087 10.547 10.479 9.618 8.490 9.724 10.411 11.810 10.158 9.375 8.017 7.094 10.148 7.900 6.608 5.682 4.562 7.083 7.486 6.052 6.052 5.203 5.337 6.339 6.339 4.643 3.810 78.477
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79.747
79 10.94 10.72 10.08 10.08 112.46 112.46 111.52 110.00 111.00 110.00 111.00 14.405 15.486 15.111 12.885 11.597 11.718 12.827 11.125 11.793 13.100 628 77.248 76.074 77.341 77.311 A SAN 10554 10554 10557 10559

7	9	9	8	9	9	9	9	7	9	9	80	9	9	9	9	9	9	9	<b>6</b> 0	7		57	89	9	9	8	7	7	9	9	8	9	9	89	7	7	9	9	80	7	9
.19	.39	.91	. 52	.56	.34	.21	. 60	69.	91	13.54	.53	12.62	.80	13.28	14.10	14.81	17.28	17.89	18.31	11.59	1.13	14.46	.41	12.78	.85	.91	23.13	11.88	12.91	.10	1.15	15.18	1.25	25.62	26.09	11.86	.63	.70	.41	.77	.28
.00 12		.00 10	.00 14	1.00 10	.00 14		.00 15		00		.00 12	.00 12			.00 14	.00 14		.00 17	00 18			.00 14	00 15	1.00 12	.00 15	.00 14		.00				.00 15	.00 28	.00 25	.00 26	.00 11	.00 15	.00	.00 16	.00 13	.00 14
543 1	365 1	-	_		1 860	294 1	447 1	_		_	_	671 1	694 1	185 1	359 1	273 1	479 1	959 1	_	_	048 1	659 1	-		386 1	460 1	9.790	455 1	056 1	619	804 1	305 1	289 1	10.579 1	7	344 1	061 1	007	994 1	209 1	220 1
6	7				12.					9.																					7.			10.	9.	æ	7.	9	4	9	ņ.
70.305	69.594	68.798		70.684	70.222	69.618	71.409	67.560	66.675	66.234	66.203	65.392	65.699	66.240	65.330	66.545	62.609	66.206	66.444	65.787	65.261	63.801	63.527	66.068	65.607	65.084	65.851	62.881	61.460	61.265	60.495	60.894	59.363	58.741	58.722	61.857	61.698	62.727	62.836	63.414	64.406
27.188	26.192	.298	.907	25.345	.344	1.067	1.436	.047	1.178	1.007	1.116	24.945	.104	7.275	25.983	3.295	27.004	28.139	29.227	21.932	.774	20.582	0.020	19.542	3.280	3.376	1.115	21.153	.160	21.628	21.059	.763	.772	20.803	18.647	.797	23.494	23.096	23.819	975	.535
2,	36	53	56	25	5	25	2	5	5	2	2	5	5	.7	2	5	7	2	2	.2	ñ	3(	5	ä	7	Ä	H	73	7	74	.21	ä	ä	7	16	5	2.	5	ci	2	2
149	149	149	149	149	149	149	149	150	150	150	150	150	150	150	1.50	150	150	150	150	151	151	151	151	151	151	151	151	152	152	152	152	152	152	152	152	153	153	153	153	154	154
LEU A	LEU A	LEU A			LEU A	LEU A	LEU A	TYR A		TYR A		TYR A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A		ASN A	ASN A	ASN A		ASN A	ASN A			GLY A		THR A	THR A							
z	F				8		CD2										CE2							g							0				~				0	z	ð
1180	1181	1182	1183	1184	1185	1186	1187	1188	1189	1190	1191	1192	1193	1194	1195	1196	1197	1198	1199	1200	1201	1202	1203	1204	1205	1206	1207	1208	1209	1210	1211	1212	1213	1214	1215	1216	1217	1218	1219	1220	1221
æ	9	œ	9	7	9	9	œ	9	9	9	9	9	9	9	7	9	9	œ	9	7	9	9	œ	9	v	9	œ	œ	7	ø	9	œ	7	9	9	œ	7	9	و	89	9
15.67	69.9	18.35	5.03	15.48	12.99	14.57	12.49	14.72	13.33	14.66	10.60	14.87	12.09	16.30	14.83	14.49	14.94	13.97	12.28	11.43	10.19	10.66	11.84	11.91	11.68	15.77	20.95	21.22	12.55	12.03	15.18	3.95	2.57	12.28	14.09	11.87	11.40	3.48	11.58	2.14	4.52
1.001	1.00.1	1.00 1	1.00 2	1.00 1	1.00.1	1.00 1	1.00.1	1.00 1	1.00 1	1.00 1	1.00 1	1.00 1	1.00 1	1.00 1	1.00 1	1.00 1	1.00 1	1.00 1	1.00 1	1.00.1	1.00 1	1.00 1	1.00 1	1.00 1	1.00 1	1.00.1	1.00 2	1.00 2	1.00 1	1.00.1	1.00 1	1.00.1	1.00 1	1.00 1	1.00 1	1.00 1	1.00.1	1.00 1	1.00 1	1.00 1	1.00 1
.961	.622	3.211	.368	.007	.285	.908	.026	.238	.546	699.	619.	878	849	975	1.264	.884	976.	.784	.323	905.1	.645	7.520	,266	1.180	1.042	1.697	9.208	1.743	7.898	.783	1.739	1.511	.790	. 703	1.267	. 903	1.258	1.707	9.729	.557	7.660
0																																									
71.840		74.288		73.874	73.314	72.384	72.837	74.481	73.93	73.45	73.90	72.946	73.38	72.91	71.17	70.25	70.13	69.87	70.683	70.13	70.06	71.386	71.51	68.94	67.57	66.467	99.99	65.351	72.533	73.82	73.97	74.880	73.157	73.265	72.415	72.644	71.583	70.942		69.390	69.927
.798	40.105	1.261	.653	.533	0.027	3.953	37.807	40.412	.905	2.192	0.054	2.677	.556	1.842	9.342	3.380	7.165	7.369	066.7	5.942	1.744	1.063	2.824	177.8	1.408	1.591	32.530	1.122	1.677	1.021	3.799	3.025	2.774	1.639	30.439	9.372	.552	29.343	28.495	9.025	29.861
38	4	4	36	35	4	36	9,1	4(	4	4	4	4	4	4	36	36	'n	'n	m	36	ň	3,	Ä	æ	Ä,	3	33	34	3,	34	æ	33	ž	3	3	55	3(	52	7	5	2
142	142	142	142	143	143	143	143	143	143	143	143	143	143	143	144	144	144	144	144	145	145	145	145	145	145	145	145	145	146	146	146	146	147	147	147	147	148	148	148	148	148
THR A	-	THR A	THR A	PHE A	PHE A	PHE A	PHE A	PHE A	PHE A	PHE A	PHE A	PHE A	PHE A	PHE A	ALA A				ALA A			GLU A				GLU A		GLU A	GLY A	GLY A	GLY A	ALA A	ALA A			ALA A					
0	-	001	-		ð		0																				-	_								_		ජ	Ü		
1138	1139	1140	1141	1142	1143	1144	1145	1146	1147	1148	1149	1150	1151	1152	1153	1154	1155	1156	1157	1158	1159	1160	1161	1162	1163	1164	1165	1166	1167	1168	1169	1170	1171	1172	1173	1174	1175	1176	1177	1178	1179



WO 99/43794

<b>9</b> 80		9 1	. 9	9	7	7	9	ø	ω	9	9	7	9	9	7	7	9	9	8	9	9	8	7	7	9	9	80	7	9	9	80	9	9	80	89	7	9	9	80
1.00 11.69	1.00 11.57	1.00 11.97	1.00 11.36	1.00 12.02	1.00 10.80	1.00 11.77			1.00 16.92	1.00 13.28	1.00 12.02	1.00 15.51	1.00 13.12	1.00 16.69							1.00 17.12			1.00 13.14	1.00 15.42	1.00 16.69		1.00 14.76		1.00 12.76	1.00 11.79	1.00 14.28					1.00 11.65	1.00 14.87	1.00 15.09
14.195	16.482	17.376	17.274	19.195	18.427	13.797	13.015	13.672	13.053	11.533	10.888	10.069	10.915	9.689	10.156	14.962	15.733	16.147	16.949	17.054	16.719	17.252	15.861	15.601	15.964	17.210	17.722	17.526	18.622	19.976	20.131	18.369	17.268	16.822	16.815	20.969	22.362	22.646	22.337
82.836	82.658	82.347	81.657	81.518	82.111	84.147	84.629	85.872	86.974				83.088	81.863	81.953	85.778		87.040				86.168			86.166	85.275	84.706	85.180	84.346	84.996	86.248	84.152	83.139	82.387	83.124	84.127	84.670	85.307	84.670
27.266	26.361	25.157	23.838	24.195	23.233	27.295	28.474	29.029	29.174	28.083	27.535	28.327	26.306	27.639	26.409	29.387	29.735	31.201	31.554	28.916	27.430	26.595	27.046	32.013	33.444	33.728	32.817	34.993	35.473	35.292	35.410	36.980	37.273	36.398	38.451	35.073	35.136	36.500	37.508
HIS A 169 HIS A 169	HIS A	HIS A		HIS A		HIS A	HIS A	HIS A	HIS A	HIS A	HIS A	HIS A	HIS A	HIS A	HIS A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	GLY A	GLY A		GLY A	ASP A	ASP A 173	ASP A	ASP A		ASP A	ASP A		Ø	ILE A 174	ILE A 174	ILE A 174
ပဝ			3 CD2	4 CE1	S NE2		7 CA					2 ND1														ပ 9								4 OD1			7 Q	8 U	0
1348	135	135	135	1354	135	135	135	135	135	136	136	136	136	136	136	136	136	136	136	137	137	137	137	137	137	1376	137	137	137	138	138	1382	1383	1384	1385	1386	1387	1388	138
œ vo	<b>&amp;</b> (	o t	۰ ۰	9	80	9	ø	9	9	7	7	9	9	80	7	9	9	œ	9	9	9	9	9	9	9	œ	7	9	9	ю	9	9	9	9	9	9	9	7	9
1.00 25.38		1.00 32.67	1.00 16.62	1.00 18.03	1.00 17.25					1.00 18.44	1.00 14.27		1.00 14.80	1.00 14.92						1.00 14.63	1.00 13.59	1.00 15.06	1.00 13.53									1.00 11.52	1.00 10.63	1.00 13.81	1.00 12.04	1.00 12.05	1.00 12.13	1.00 11.43	1.00 12.52
5.002	2.496	1.658	6.929	8.391	9.173	6.826	5.423	5.334	5.245	4.349	8.700	10.082	11.109	12.287	10.734	11.732	12.739	13.776	11.022	10.103	10.291	9.121	9.421	8.254	8.469	7.594	12.342	13.244	13.551	12.854	12.493	12.174	13.130	10.899	12.765	10.574	11.518	14.709	14.979
80.605	80.146	82.359	79.430	79.888	79.767	78.140	77.497	76.186	76.297	75.308	80.499	81.024	79.863	80.096	78.765	77.692	78.009	77.349	76.374	75.836	76.232	74.921	75.722	74.385	74.809	74.289	78.870	79.208	80.677	81.539	78.934	77.459	76.583	77.007	75.214	75.639	74.751	81.067	82.473
18.055			20.086		18.735		20.215	•	•	23.067	•		•		•							•				18.235						26.552		•			26.752	•	25.979
O THR A 164 CB THR A 164	THRA	THRA	LYS A	LYS A	LYS A	CB LYS A	CG LYS A	CD LYS A	CE LYS A	NZ LYS A	N GLY A	CA GLY A	C GLY A	O GLY A	N TYR A	CA TYR A	C TYR A	O TYR A	CB TYR A	CG TYR A	CD1 TYR A	CD2 TYR A	CE1 TYR A	CE2 TYR A	CZ TYR A	OH TYR A	N PHE A	CA PHE A	C PHE A	O PHE A	CB PHE A	CG PHE A	CD1 PHE A	CD2 PHE A	CE1 PHE A	CE2 PHE A	CZ PHE A	N HIS A	CA HIS A
1306	1308	309	1311	312	313	314	315	316	317	318	315	32(	32.	32	323	324	325	326	327	328	323	33	331	332	33	334	33	33(	33,	338	33	340	343	342	343	344	345	346	347

9 8	w	9	œ (	۰ ر	9	9	80	9	9	9	۲	φ	- 1		. 19	9	8	9	0	، و	۰ و	o u		<b>&amp;</b>	7	ø	9 0	20 \	۰ ب	م	9	89	<b>a</b> o	7	9	9	ထ
1.00 12.33	15	18	1.00 20.56		11						11.3			1.00 13.64								1.00 18.03					10.4								1.00 11.02	1.00 11.67	1.00 10.91
28.903	28.184	27.763	28.431	29.477	29.205	27.757	27.225	30.172	31.654	31.899	31.571	30.812	30.359	30.389	25.688	24.870	24.044	25.196	23.698	23.179	22.837	21.808	20.988	19.614	25.083	24.265	24.504	23.582		24.214		23.958	24.690	25.796	26.110	25.471	25.197
87.603	89.341	90.225	91.232	87.768	86.771	86.833	85.746	87.051	86.796	85.340	84.389	83.305	82.893	82.28 87.990	87.980	87.277	86.403	89.446	89.417	89.009	89.736	88.936	89.265	89.156	87.528	86.836	85.311	84.533	87.266	88.769	.04	88.229	90.199	84.945	83.521	82.709	81.542
30.292	31.997		30.462	29.115	28.057	27.585	27.261	26.893	27.286	27.797	26.694	26.896	28.090	27.75	27.104	28.195	27.826	26.915	26.645	25.446	27.712	25.242	26.275	26.097	29.473	30.468	30.442	30.482	31.939	32.131	33.640	34.487	34.009	30.314	30.436	29.302	29.555
ASP A 179 ASP A 179	ASP A	ASP A	4	ARG A 180	ARG A	ARG A	ARG A	ARG A	ARG A	ARG A	ARG A	ARG A	ARG A	NH2 AKG A 180	TYR A	TYR A	TYR A	TYR	TYR A	TYR A	<b>4</b>	CEL TYR A 181	TYR A	TYR A	GLU A	GLU A	<b>A</b> ,	GLU A	GLU A	GLU A	GLU A	GLU A	OE2 GLU A 182	ALA A 183	ALA A	ALA A 183	ALA A
1432 C				1437 U						_				1448 N						1455 C		1457 C											1469 0	1470 N		1472 C	
99	9	9	۲,	و ه	ω.	9	80	7	9	9	<b>a</b> o	9	<b>6</b>	ין מ		9	9	æ	9	9	<u>ب</u> م	م و	· vo	9	9	ø	91	- (	۰ م	9	œ	9	9	ထ	80	7	9
3.27	4	7	. 22	. 63	20	98.	24.28	14.39	13.41	3.76	.43	.61	.91	15.18	14.29	13.39	15.05	14.25	16.12	14.43	16.58	13.92	18.68	17.34	14.28	17.20	18.00	12.43	11.08	12.88	15.73	17.78	4.33	18.07	24.77	11.79	11.42
1.00 1		11	14	1.00 18.										1.00 23								1.00.1		1.00 1								1.00 17	1.00 24	1.00 1	1.00	1.00	1.00
23.357 1.00 1:	806 1.00	000 1.00	1.00 14	770 1.00	938 1.00	138 1.00	1.00		1.00		1.00	1.00	243 1.00	0 0		1.00	1.00	1.00	1.00	1.00	1.00		1.00	1.00	1.00	1.00	1.00		1.00	1.00	.881 1.00	1.00	.362 1.00		33.777 1.00	29.810 1.00	29.357 1.00
.357 1.00	24.806 1.00	24.000 1.00	23.260 1.00 14	25.291 1.00	25.938 1.00	23.138 1.00	23.724 1.00	25.854 1.00	27.336 1.00	27.678 1.00	27.482 1.00	27.724 1.00	29.243 1.00	0 0	28.237 1.00	28.558 1.00	29.724 1.00	29.909 1.00	28.953 1.00	27.721 1.00	26.998 1.00	1.00	26.014 1.00	27.392 1.00	25.169 1.00	26.574 1.00	25.441 1.00	30.469 1.00	5 31.517 1.00	31.053 1.00	31.881 1.00	.717 32.739 1.00	33.362 1.00	.440 1.00	155 33.777 1	.608 29.	88.502 29.357 1.00
23.357 1.00	84.016 24.806 1.00	81.690 24.000 1.00	86.493 23.260 1.00 14	87.131 25.291 1.00	86.587 25.938 1.00	88.470 23.138 1.00	89.044 23.724 1.00	87.662 25.854 1.00	87.678 27.336 1.00	86.561 27.678 1.00	86.760 27.482 1.00	89.032 27.724 1.00	89.123 29.243 1.00	88.166 29.982 1.00	85.465 28.237 1.00	84.361 28.558 1.00	84.677 29.724 1.00	83.898 29.909 1.00	83.145 28.953 1.00	82.559 27.721 1.00	83.101 26.998 1.00	81.291 27.142 1.00	81.131 26.014 1.00	80.298 27.392 1.00	80.031 25.169 1.00	79.182 26.574 1.00	79.045 25.441 1.00	85.795 30.469 1.00	86.126 31.517 1.00	87.115 31.053 1.00	87.420 31.881 1.00	86.717 32.739 1.00	85.739 33.362 1.00	84.535 33.440 1.00	86.155 33.777 1	27 87.608 29.	48 88.502 29.
ILE A 174 34.896 83.495 23.357 1.00	CG2 ILE A 174 35.145 84.016 24.806 1.00	CD1 ILE A 174 33.220 81.690 24.000 1.00	A 175 36.441 86.493 23.260 1.00 14	CA SER A 1/5 37.710 87.093 23.770 1.00 C SER A 175 37.712 87.131 25.291 1.00	O SER A 175 38.617 86.587 25.938 1.00	CB SER A 175 37.868 88.470 23.138 1.00	OG SER A 175 39.049 89.044 23.724 1.00	N ASN A 176 36.650 87.662 25.854 1.00	CA ASN A 176 36.515 87.678 27.336 1.00	C ASN A 176 35.511 86.561 27.678 1.00	O ASN A 176 34.286 86.760 27.482 1.00	CB ASN A 176 35.898 89.032 27.724 1.00	CG ASN A 176 35.749 89.123 29.243 1.00	ODI ASN A 176 35.963 88.166 29.982 1.00	N TRP A 177 36.085 85.465 28.237 1.00	CA TRP A 177 35.172 84.361 28.558 1.00	C TRP A 177 34.248 84.677 29.724 1.00	O TRP A 177 33.279 83.898 29.909 1.00	CB TRP A 177 36.054 83.145 28.953 1.00	CG TRP A 177 36.712 82.559 27.721 1.00	CD1 TRP A 177 37.745 83.101 26.998 1.00	TRP A 177 36.399 81.291 27.142 1.00 mpp a 177 38 070 82 235 25 940 1 00	CE2 TRP A 177 37.234 81.131 26.014 1.00	CE3 TRP A 177 35.437 80.298 27.392 1.00	CZ2 TRP A 177 37.148 80.031 25.169 1.00	TRP A 177 35.379 79.182 26.574 1.00	CH2 TRP A 177 36.253 79.045 25.441 1.00	N ASP A 178 34.477 85.795 30.469 1.00	ASP A 178 33.507 86.126 31.517 1.00	C ASP A 178 32.454 87.115 31.053 1.00	O ASP A 178 31.586 87.420 31.881 1.00	CB ASP A 178 34.243 86.717 32.739 1.00	CG ASP A 178 35.201 85.739 33.362 1.00	OD1 ASP A 178 34.916 84.535 33.440 1.00	ASP A 178 36.317 86.155 33.777 1	N ASP A 179 32.527 87.608 29.	CA ASP A 179 31.448 88.502 29.

15.82 19.99 15.14 12.41 22.589 20.788 22.180 20.136 24.204 25.327 25.350 21.504 20.769 20.639 21.186 20.929 19.861 20.365 20.781 20.943 20.956 21.885 21.756 20.648 20.648 23.067 78.658 778.457 776.558 776.558 776.559 776.589 776.989 777.103 34.321 35.600 34.131.682 33.682 33.364 35.865 33.364 35.865 35.081 36.737 38.000 38.851 38.000 38.851 39.066 38.920 38.777 38.777 38.777 38.777 47.403 47.403 44.691 44.691 44.691 44.691 43.951 42.769 44.778 45.446 45.279 46.014 1541 1542 1543 1544 1545 1546 1546 10.70 8.62 10.69 10.72 12.35 9.80 9.80 12.56 15.21 8.78 11.90 9.74 12.61 10.95 12.59 12.07 14.15 13.43 13.43 14.75 11.31 11.31 11.31 11.31 11.09 1.00 1.00 1.00 1.00 1.00 21.805 23.587 22.703 24.494 20.243 20.601 20.286 19.726 19.379 20.965 035 20.127 20.238 19.195 20.314 19.065 22.226 20.428 981 84.887 85.003 85.530 83.352 83.381 82.624 82.844 81.913 83.078 82.584 83.315 84.584 84.608 84.298 84.298 84.846 85.361 85.004 84.254 84.170 85.647 85.390 85.166 85.345 86.646 87.655 224 943 896 30.290 28.196 26.939 26.939 26.939 26.939 27.046 28.25 27.186 27.186 28.25 27.186 28.25 27.186 28.25 28.25 29.469 28.25 29.69 20.69 20.71 20.7 SELAN 

10.61 11.55 11.79 11.94 11.94 11.94 11.60 9.83 11.55 1 11.04 10.64 19.700 21.151 21.208 18.862 18.667 17.277 16.903 18.875 20.252 20.330 21.434 16.435 15.071 16.505 16.950 17.499 16.249 18.837 19.473 19.371 15.372 16.008 74.135 72.879 74.045 79.160 79.160 77.91 77.91 77.91 78.294 77.68 81.011 78.294 77.359 77.359 76.330 76.231 77.98 78.68 78.68 77.359 77.359 76.231 78.298 78.298 80.71 78.298 78.298 80.731 81.166 82.268 82.268 82.268 82.268 82.268 83.276 84.276 83.276 84.276 84.276 86.776 87.276 88.7776 88.776 88.776 88.776 88.776 88.776 88.776 88.776 88.776 88.7776 88.776 88.776 88.776 88.776 88.776 88.776 88.776 88.776 88.7776 88.776 88.776 88.776 88.776 88.776 88.776 88.776 88.776 88.7776 88.776 77.124 79.108 78.219 78.098 77.148 78.892 78.307 77.115 22.499 23.465 118.590 117.434 16.849 15.856 17.877 18.522 18.421 17.305 19.852 20.002 20.680 20.457 17.507 16.939 18.105 17.604 25.178 24.799 24.658 23.024 22.055 20.810 19.994 21.636 20.738 20.786 21.042 22.393 16.020 16.441 17.271 17.611 14.797 14.802 891 1609 1614 1616 1617 1618 1619 1620 1622 1623 1624 1625 1626 1612 1613 1627 1628 1630 1631 633 1629 11. 84 11. 03 11. 03 11. 69 11. 46 11. 46 11. 26 12. 24 12. 03 10. 73 11. 52 12. 52 12. 52 12. 52 13. 52 14. 52 15. 52 16. 52 17 9.97 10.64 14.16 11.22 10.28 10.50 11.99 13.33 12.29 10.72 11.33 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 16.536 17.750 14.855 15.767 16.649 15.747 17.460 17.450 15.895 16.704 16.423 15.519 16.186 16.216 17.262 17.060 16.584 18.410 20.334 14.905 15.830 13.565 16.236 17.900 17.112 18.134 16.795 19.353 19.671 19.107 .481 .314 72.942 72.533 72.181 77.497 77.782 76.649 75.902 77.416 74.883 74.298 73.454 76.637 76.658 75.622 75.928 80.119 76.594 73.384 75.877 77.869 77.880 76.976 77.265 77.423 75.629 74.412 74.063 79.059 75.622 79.433 79.529 80.452 76.847 76.690 34.007 32.756 32.364 33.660 34.880 34.439 35.719 30.995 29.362 29.429 28.459 27.030 26.607 28.744 28.236 28.236 40.612 39.441 39.621 40.411 40.568 41.707 39.688 41.871 40.860 38.283 37.097 36.081 36.284 36.416 37.442 35.060 27.401 444 

14.64 10.02 10.72 10.18 11.96 10.83 12.55 10.81 12.47 10.73 12.42 12.22 12.33 15.18 13.61 12.75 10.21 10.82 11.60 13.68 15.03 14.89 19.713 18.893 19.480 18.920 20.069 20.933 22.677 22.875 22.302 22.903 22.283 20.496 21.335 20.889 21.373 21.808 19.170 18.671 18.997 22.234 21.193 21.577 65.829 66.477 66.643 66.828 66.828 65.326 64.422 64.023 62.882 62.882 64.023 64.023 64.970 64.970 62.667 65.876 63.854 62.923 61.577 60.552 63.523 64.740 70.688
71.237
72.628
70.260
69.327
68.689
68.689
69.327
60.354
69.324
69.327
60.354
69.327
60.842
60.842
60.842
60.842 61.496 19.388 20.603 20.603 21.766 116.539 116.539 114.193 114.488 13.33 114.813 116.998 116.998 116.998 116.998 116.748 116.748 116.748 116.748 116.748 116.748 116.748 18.422 18.819 19.679 18.155 18.185 17.073 15.885 14.724 15.035 14.673 12.285 12.982 15.759 16.153 16.16364 13.100 18.376 17.246 18.038 1685 1685 1686 1687 1689 1690 1691 1693 1693 1693 1696 1698 1698 1699 1700 1702 1703 1704 1705 1706 1709 1709 14.40 18.35 10.31 10.54 10.54 10.61 9.16 112.29 111.91 112.96 113.37 113.37 114.22 114.22 117.39 117 11.12 11.54 12.56 11.38 12.87 16.27 13.55 13.50 14.21 10.88 10.76 11.55 1.000 1.00 1.00 1.00 16.979 17.231 16.273 13.218 17.021 18.339 18.311 18.883 17.236 16.877 17.426 16.136 16.197 18.180 16.062 15.724 14.743 14.457 13.584 15.060 13.286 13.915 11.497 10.662 10.930 14.188 15.081 16.457 17.051 15.212 13.844 17.957 17.821 027 72.638 75.017 75.155 74.624 74.624 74.562 74.185 72.053 75.359 71.431 69.200 71.573 70.429 69.230 69.372 67.727 69.989 69.148 68.083 69.485 67.341 68.739 67.676 66.968 72.621 74.603 75.060 73.256 74.586 73.996 11.371 70.454 58.930 69.461 11.847 11.089 10.565 11.168 16.161 15.866 14.760 16.344 17.295 18.330 18.330 19.481 20.193 20.550 20.691 16.912 16.912 16.912 11.714 11 15.876 17.840 19.072 19.031 20.269 20.184 17.634 21.368 22.527 17.985 18.731 17.570 21.432 **AAAAAAAAAAA** 

64	
	000000000000000000000000000000000000000
11.67 10.83 11.68 11.68 11.67 11.67 11.60	10.94 10.94 10.94 11.29 11.29 10.28 11.21 11.21 15.31 9.30 9.30
30.709 31.951 32.306 33.356 33.356 33.356 33.123 33.926 30.106 30.83 30.907 30.907 31.228 30.178 30.178 30.	31.114 31.114 31.114 29.126 29.218 29.269 29.378 20.378 20.378 20.412 20.412 20.412 20.412 20.412
	68 68 79 40 68 90 1 69 75 1 70 45 0 69 78 3 68 8 14 70 60 1 70 60 1 70 69 67 2 70 39 9 71 71 915
19.009 18.391 18.621 18.116 18.997 19.610 19.610 19.186 19.393 20.641 21.062 22.36 2	24.531 24.533 25.807 26.042 23.295 22.096 23.196 20.743 26.544 27.846
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	24. 24. 24. 25. 23. 23. 20. 20. 26.
	10000000000000000000000000000000000000
7	
ASP ASP ASP ASP ASP ASP ASP GUY GUY GUY LEU LEU LEU LEU LEU LEU LEU LEU LEU LEU	ARG 11LE 11LE 11LE 11LE 11LE 11LE 11LE 11L
Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z Z	NH2 N C C C C C C C C C C C C C C C C C C C
1768 1770 1771 1772 1773 1774 1775 1776 1778 1781 1782 1788 1788 1788 1789 1799 1790 1790 1791 1792 1793 1793	798 800 801 803 803 805 805 805 806
	00000000000000000000000000000000000000
18.32 21.98 110.07 111.16 111.16 111.16 112.00 113.19 114.29 114.29 114.29 114.29 114.29 114.29 114.29 114.29 114.29 114.29 114.29 114.29 115.25 116.64 117.29 117.	16.42 16.42 17.85 111.47 111.21 13.22 13.24 14.03 14.03 13.32 14.03
17.261 16.335 22.269 23.261 24.754 25.269 24.754 25.3876 25.3876 25.378 25.378 26.376 27.107	20.077 18.823 18.734 25.655 26.999 27.965 27.426 27.426 28.356 29.53 29.653 29.653
59.162 59.976 60.043 59.317 59.317 59.317 59.317 60.052 60.053 60	55.624 53.916 54.666 54.129 55.060 54.635 56.111 56.912 57.794
17.856 18.762 17.392 17.392 17.396 18.781 17.885 19.887 19.887 11.9887 11.993 11.993 11.914 11.31 11.3	928 928 928 964 965 2297 139 138
19.76   19.76	9.7.89.7.66.00
	. — — — — — — — — — — — — — — — — — — —
220 220 220 220 220 220 220 220 220 220	220 220 220 221 221 221 221 222 222 222
	A 220 A 220 A 220 A 221 A 221 A 222 A 222 A 222 A 222 A 222
	HIS A 220 HIS A 220 HIS A 220 GUY A 221 GUY A 221 GLY A 221 ALA A 222
GLIN A GL	CD2 HIS A 220 CE1 HIS A 220 NE2 HIS A 220 N GLY A 221 C GLY A 221 O GLY A 221 N ALA A 222 C ALA A 222

65 11.83 12.34 8.31 9.45 9.94 10.71 12.93 12.50 10.79 10.78 8.67 10.81 10.03 7.85 10.81 11.07 10.75 9.60 10.90 1.000 25.369 24.740 25.519 23.448 29.396 29.759 31.098 30.914 28.312 27.882 27.289 27.552 26.370 25.702 29.177 26.526 24.467 23.691 22.358 75.588 76.331 74.503 77.219 74.340 80.148 80.799 80.799 80.859 80.80 80.590 80.590 80.590 80.590 80.590 78.402 80.196 80.269 80.067 81.403 80.140 79.857 79.857 77.638 76.864 75.738 74.641 75.595 75.242 74.740 74.046 73.200 75.939 74.844 74.489 775.303 775.303 24.335 22.023 23.982 22.023 22.629 22.023 22.036 20.367 20.367 20.132 20.036 18.981 18.975 19.739 19.596 19 16.476 17.642 16.442 18.751 16.367 18.634 17.468 17.965 18.119 PHE A A SW A A SW 1865 1865 1867 1867 1869 1870 1870 1871 1872 1873 880 881 882 883 1876 984 9 . 88 9 . 23 110 . 24 110 . 28 9 . 93 9 . 73 9 . 90 9 . 9 9.37 12.72 12.48 9.62 9.94 9.35 11.03 11.85 9.67 10.57 9.21 11.50 9.50 27.464 26.841 29.487 29.548 29.753 30.634 32.047 30.234 28.883 28.809 31.209 30.602 31.758 32.236 33.104 27.814 26.480 25.537 24.388 25.915 28.049 25.941 25.062 25.991 26.769 26.769 26.587 27.505 25.574 28.582 28.257 28.257 26.674 27.513 72.338 69.865 70.442 70.228 71.128 72.633 74.068 74.868 75.695 74.180 74.708 76.905 74.836 74.934 78.763 79.391 80.604 79.043 78.801 78.817 80.157 78.589 79.044 78.129 78.345 79.598 77.530 79.483 78.757 80.205 26.757
28.521
29.904
30.725
30.726
30.726
30.726
28.575
27.456
27.456
28.727
28.727
28.727
28.728
28.728
28.729
28.7338
28.7338 30.937 32.215 27.364 27.744 26.576 26.698 28.849 30.145 30.780 31.838 25.342 24.165 23.015 ASP A 228
ASP A 228
ASP A 228
ASP A 228
ALA A 229
ALA A 229
ALA A 229
ALA A 229
VAL A 230
VAL A 231
VAL A 232
VAL A 232 

66 11.59 13.03 12.53 9.85 9.85 11.79 11.71 11.60 11.60 11.49 11.60 11 30.770 31.500 32.359 32.995 33.408 33.408 34.021 34.208 34.816 29.672 24.483 24.133 25.269 27.593 28.039 29.205 29.219 28.461 27.206 27.206 27.692 26.282 30.136 31.258 29.112
28.581
28.651
27.940
28.518
27.340
28.7139
27.367
28.7139
28.7139
28.7139
28.7139
28.7139
28.7139
28.7139
28.7139
28.7139
28.7139
28.7139
28.7139
28.7139 70.400 71.869 71.869 72.780 65.066 65.200 66.200 66.200 66.200 66.037 66.037 66.037 66.030 9.398 9.129 9.129 112.2213 112.730 111.655 111.665 111.665 111.665 111.665 111.665 111.665 111.665 111.665 111.665 111.665 111.928 111.928 111.928 111.928 112.163 113.319 113.319 114.103 115.227 117.665 117.665 117.665 117.665 117.665 117.665 117.665 118.665 119 [934 [937] [938] [ 10.06 10.191 11.91 11.91 11.91 11.11 11.11 11.12 11.13 29.757
30.433
31.183
31.182
33.246
33.246
34.246
26.736
26.736
27.330
26.736
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330
27.330 .484 .998 .763 .122 75.523 77.568 74.684 77.186 76.598 77.442 77.442 77.644 77.644 77.644 77.644 77.644 77.644 77.644 77.64 77.64 77.65 77.6 69.448 68.474 71.904 71.717 71.113 72.178 69.536 68.449 67.097 66.061 68.728 70.142 

9.94 111.26 10.35 111.23 111.23 111.23 111.03 111.03 111.23 111.23 111.23 111.23 111.23 111.23 111.3 34.913 35.914 33.912 31.562 31.562 31.662 32.016 33.921 33.921 33.921 33.921 33.921 33.921 33.921 34.961 33.921 34.961 34 34.412 67.083 67.647 67.378 66.766 67.378 68.001 68.0179 68.0179 68.0179 68.131 70.087 71.1308 71.17 20.498 20.094 118.694 117.749 117.749 119.694 110.608 111.761 122.750 123.707 123.707 123.707 123.707 123.707 124.552 125.608 126.095 127.708 128.109 128.109 129.1 LEU A
LEEU A
LEEU A
LEEU A
LEEU A
LEEU A
VAL A
COLU A
COLU
COLU A
COLU
COLU A
COLU
COLU A
CO 22. 20 29. 85 118. 09 21. 86 11. 94 11. 94 11. 94 11. 94 11. 94 11. 94 11. 94 11. 95 11. 96 1 34.494 34.626 35.728 35.728 35.030 35.030 33.030 33.593 34.467 34.990 31.503 34.467 34.990 31.503 34.467 34.900 34 . 292 . 977 . 150 . 885 . 830 . 917 . 984 . 876 . 542 . 542 . 543 . 543 . 543 . 543 . 543 . 543 . 543 . 543 . 543 . 543 . 543 . 543 . 543 . 644 . 645 . 645 . 645 . 646 . 646 . 646 . 646 . 646 . 647 288 60.676 60.755 59.497 58.367 61.400 61.400 61.213 62.316 52.316 52.316 57.175 54.836 60.678 61.336 60.354 60.354 60.891 59.405 59.027 61.541 61.346 62.413 63.303 61.331 60.064 63.784 62.248 63.182 64.105 63.587 63.290 63.951 63.494 64.802 64.327 65.031 65.389 66.278 10.846 11.7551 10.7779 10.7379 10.7379 10.7379 11.5520 

68 16.68 14.50 115.71 111.26 111.25 111.35 111.63 112.23 113.10 113.10 113.10 113.10 113.10 113.10 113.10 113.10 113.11 113. 81 ë. 40.988 38.110 34.822 34.146 39.353 40.271 40.469 41.616 43.640 39.957 38.316 37.793 37.837 37.199 37.632 35.855 42.410 43.375 40.726 41.471 41.270 40.599 41.019 40.377 40.214 40.998 39.536 38.957 34.889 86.990 86.447 87.990 88.396 87.407 88.302 87.901 88.302 87.901 88.305 84.455 83.997 83.268 85.889 86.052 85.151 87.536 32.579 31.378 33.134 34.378 35.185 34.455 34.455 36.698 31.684 31 2097 2098 2099 2100 2100 2102 2102 2104 2107 2109 2110 2111 21113 12.06 13.57 111.31 111.37 113.71 113.71 113.38 116.44 20.59 9.86 112.24 112.24 113.01 113.01 110.69 110.69 110.69 110.69 111.65 111.66 111.73 1.00 1.00 34.631 36.550 37.361 37.891 38.227 37.891 38.486 39.947 40.317 37.665 36.269 36.269 35.274 40.708 42.158 42.936 42.502 42.074 41.181 42.569 44.230 35.355 34.869 33.335 33.723 34.524 33.234 34.859 33.564 033 078 445 938 690 170 091 ž. **4**6. £6. 5 76.859 77.988 75.508 74.978 75.687 75.687 73.130 73.206 73.206 73.306 73.306 73.908 73.908 73.908 73.909 74.909 76.605 77.705 77 80.239 82.528 78.518 77.881 78.035 81.124 82.197 83.428 81.286 81.807 82.593 81.502 80.528 83.254 81.583 086 856 35.642 31.751 31.155 32.085 33.325 30.973 30.818 32.527 32.370 33.189 33.189 33.189 33.189 33.199 33.707 33.030 33.584 33.584 33.586 33 30.818 31.873 31.385 32.178 

11. 83 11. 31 11. 31 11. 97 12. 55 12. 25 12. 25 14. 18 9 94 14. 75 16. 83 16. 12 10. 52 11. 16 11. 31.956 38.308 39.008 39.008 39.008 40.039 40.092 40.098 40.098 40.098 42.004 43.107 43.214 43.214 43.214 38.32 36.165 37.632 74.258
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001
77.001 21.365 20.678 20.678 19.062 18.478 19.062 19.814 19.362 20.628 18.429 20.053 20.918 22.294 11. 45 12. 17 12. 17 12. 17 10. 99 10. 09 10. 09 11. 50 11 33.3.6.4 33.3.6.4 33.3.6.4 33.3.6.4 33.6.4 33.6.6.4 33.6.6.4 33.6.6.4 33.6.6.4 33.6.6.4 33.6.6.4 83.803 84.417 85.369 81.906 81.608 81.608 81.608 81.608 81.608 81.608 81.618 81 23.490
24.122
24.906
23.720
24.906
23.720
24.678
23.079
26.221
26.221
27.725
28.236
27.725
28.236
28.236
29.600
29.600
29.600
29.600
20.725
20.847
20.883
20.983
20.983
20.983 2139 21441 2

																					7	0										•									
89	9	9	80	89	7	9	9	80	9	9	9	9	7	9	9	æ	9	9	80	7	7	9	9	œ	9	89	9	7	9	9	œ	9	9	9	7	9	9	80	9	9	9
1.00 11.00	1.00 9.55	1.00 10.92	1.00 12.04	1.00 12.59	1.00 10.30	1.00 9.08	1.00 9.26	1.00 11.04				1.00 13.68													1.00 15.82			1.00 8.64		1.00 11.04		1.00 11.02		1.00 11.61	1.00 9.86	1.00 8.30	1.00 12.22	1.00 10.89	1.00 9.93	1.00 11.46	1.00 11.35
45.468	44.545	43.692	42.710	44.043	45.128	46.061	45.505	46.238	46.481	47.790	49.023	47.999	44.168	43.586	43.872	44.184	42.039	41.392	40.654	41.597	43.719	43.907	45.292	45.486	43.745	42.397	43.984	46.268	47.655	47.985	48.642	48.605	50.051	48.191	47.430	47.642	47.144	47.806	46.851	47.591	46.690
75.092	76.068	77.042	77.631	77.214	73.464	72.569	71.940	71.757	71.419	70.697	71.580	69.510	71.683	71.091	71.937	71.401	71.000	70.507	71.318	69.245	73.275	74.106	73.906	73.741	75.596	75.753	76.459	73.847	73.732	72.341	72.184	74.237	74.258	75.632	71.287	69.911	69.830	69.251	68.979	68.833	67.579
33.806	30.773	29.979	30.499	28.773	32.248	32.959	34.219	35.179	31.968	32.367	32.357	31.384	34.226	35.441	36.687	37.751	35.269	36.571	37.171	36.866	36.588	37.769	38.367	39.593	37.365	36.906	38,641	37.440	37.915	38.401	39.438	36.780	37.297	36.323	37.764	38.262	39.715	40.554	37.342	36.004	38.025
ASP A 285	ASP A	ASP A	ASP A	ASP A	LEU A	LEU A	LEU A	LEU A	LEU A	LEU A	LEU A	LEU A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	THR A	THR A	THR A	THR A	THR A	THR A	THR A	VAL A	VAL A	VAL A	VAL A	VAL A	VAL A	VAL A	ILE A	ILE A	ILE A	ILE A	ILE A	ILE A	ILE A
0												CDS																z									ບ	0	83	CG1	ប្ជ
2265	2266	2267	2268	2269	2270	2271	2272	2273	2274	2275	2276	2277	2278	2279	2280	2281	2282	2283	2284	2285	2286	2287	2288	2289	2290	2291	2292	2293	2294	2295	2296	2297	2298	2299	2300	2301	2302	2303	2304	2305	2306
80	9	9	æ	7	7	9	9	œ	9	9	9	7	9	9	œ	9	9	9	9	7	9	9	æ	9	9	80	8	7	9	9	80	9	9	9	9	9	9	9	7	9	9
1.00 12.57			_		1.00 11.45	1.00 9.71						1.00 9.31									1.00 11.76				1.00 11.60			1.00 10.41		1.00 11.21		1.00 10.51			1.00 11.57		1.00 11.45	1.00 9.72	1.00 9.13		1.00 10.95
36.365	38.864	39.263	39.990	38.787	38.492	37.977	38.660	39.799	38.187	37.397	39.638	37.904	38.460	39.476	39.298	37.283	36.538	35.102	37.196	40.581	41.764	41.696	42.316	43.014	44.271	44.259	45.290	40.834	40.731	41.966	42.313	39.517	38.227	37.520	37.765	36.386	36.596	35.903	42.687	43.882	44.885
71.116					71.312							70.675									71.304							72.815				74.409			73.416				74.753	75.	
22.506	20.968	19.492	18.839	18.974	23.262	24.622	25.671	25.581	25.013	24.019	25.038	26.786	28.031	28.631	28.577	29.022	28.650	29.159	29.314	29.128	29.503	30.935	31.877	29.212	29.282	29.655	28.866	31.108	32.439	32.746	33.941	32.509	32.750	31.725	34.073	31.925	34.292	33.234	31.729	32.061	32.799
23 O ASN A 280	CB ASN A	CG ASN A	OD1 ASN A	ND2 ASN A	N VAL A	CA VAL A	C VAL A	O VAL A	CB VAL A	CG1 VAL A	CG2 VAL A	N LEU A	CA LEU A	C LEU A	O LEU A	CB LEU A	CG LEU A	CD1 LEU A	CD2 LEU A	N ASP A	CA ASP A	C ASP A	O ASP A	CB ASP A	CG ASP A	OD1 ASP A	OD2 ASP A	N PHE A	CA PHE A	C PHE A	O PHE A	CB PHE A	CG PHE A	CD1 PHE A	CD2 PHE A	CE1 PHE A	CE2 PHE A	CZ PHE A	N ASP A	CA ASP A	C ASP A
2223	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22	22

~		~	١.	~		_			~		10		٠.			٠.	_		١,	<u></u>	1	~	10	~	10		~	٠.		٠.	~	~	~			_	,,	_		~
,- (	, w	α)	5	<u>.</u>				~		~	~	~	~					٠.		٠.	~ ~				9		۵.	9	9	<u>پ</u>	٠.			9	9		9	ω,	9	-
10.00	13.50	12.22	11.6	15.42	13.16	11.24	12.10	17.18	17.61	11.80	14.38	13.09	11.70	12.95	12.92	11.87	13.51	13.52	13.29	15.14	13.68	12.9	15.45	10.17	13.00	14.72	14.92	11.79	14.80	16.23	15.11	17.75	13.71	11.61	14.08	11.91	15.34	15.70	16.98	11.86
000	8		00	8		00		00.	00:	00.1		80.		00.1		00.1		00.1				00	8	00.		00	00.	00	00.	8	00.	8	00	00	80	00.	80.	00.	00.	00
	1 6	14 1	3 1	-	-	~	_	_	_	-	-	_	_	_	-		٠.		_	_		_	1 09	Н	_	_	1 1	-	-	1 89	Н	7	٦	7	5 1	4 1	Н.	1 4	4	E.
46.990	48.549	48.874	46.25	46.812	44.770	49.472	50.88	50.984	51.952	51.610	52.199	51.385	53.565	51.854	54.064	53.231	50.183	50.27	50,925	51.220	48.870	48.03	48.250	51.194	51.975	53.114	53.094	51.214	50.299	49.668	48.827	50.107	54.106	55.252	55.355	54.644	56.601	56.80	56.46	56.44
		36	47	92:						171	124	11	11,	95	44	20										38	53	801	172				20		85		20	.771		672
70.499	71.89	72.436	72.647	73.276	72.308	71.6	72.0	73.543	74.035	71.2	69.924	68.	69.	67.6	68.444	67.4	74.322	75.773	76.3	77.	76.4	75.9	76.059	75.4	75.785	74.	73.753	75.808	77.072	77.320	76.525	78.376	75.020	74.106	73.485	73.874	74.850	75.7	•	•
46.271	47.675	48.755	47.162	46.005	46.976	46.760	917	47.104	47.719	48.052	47.630	47.153	47.760	821	47.438	948	46.411	398	131	45.018	488	446	47.869	44.162	43.009	42.852	43.624	41.654	41.692	40.301	39.959	39.635	42.031	41.924	518	39.580	42.152	116	43.511	337
46.	47.	48.	47.	46.	46.	46.	46.	47.	47.	48.	47.	47.	47.	46.	47.	46.	46.	46.	45.	45.	46.	45.	47.	44.	43.	42.	43.	41.	41.	40.	39.	39.	42.	41.	40.	39.	42.	41.	43.	40.
									_		_			_	_												_													
A 296	4 296	4 296	4 296	4 296	¥ 296	4 297	4 297	A 297	A 297	A 297	A 297	4 297	4 297	A 297	A 297	4 297	4 296	4 298	4 298	4 298	4 298	4 296	₹ 298	A 299	4 299	4 299	4 299	4 299	₹ 299	4 299	4 299	4 299	300	300	300	300	300	300	300	A 301
THR /	THR /	THR /	THR 1	THR 2	THR 1	PHE 1	PHE 1	PHE 1	PHE 1	PHE 1	THR /	THR /	THR 1	THR 1	THR 1	THR 1	THR 1	GEN 7	GLN 7	GLN 7	GLN 2	THR 1	MET A																	
z ő	ט	0	8	003	CG2	z	ర్	ບ	0	B	g	<del>1</del> 00	CD2	Œ1	CE2	Z	z	S S		0						ັ	0			8	OE1	NEZ	z	e G	ັບ	0	8	001	CGS	z
2349				2354	2355	2356	2357	2358	2359	2360	2361	2362	2363	2364	2365	2366	2367	2368	369	370	2371	372	373	2374	2375	376	2377	2378	2379	2380	2381	2382	2383	2384	2385	2386	2387	2388	2389	2390
NN	7	7	2	2	2	7	7	7	7	7	7	2	7	~	7	7	7	7	73	7	2	2	7	71	7	8	7	7	7	7	7	2	2	7	7	7	7	7	7	73
9 1	· w	9	80	9	9	9	7	9	7	7	7	9	ø	æ	9	9	æ	7	7	9	9	80	ø	φ	9	7	9	9	80	9	9	9	9	9	9	9	7	9	ø	œ
1.71	.67	9.59	3.39	10.96	11.39	11.11	11.15	12.12	11.32	12.38	5.65	11.32	11.68	11.50	11.01	14.57	9.84	18.55	0.40	).24	11.21	11.98	10.95	12.97	13.88	11.12	1.38	11.51	11.86	11.70	11.57	11.48	13.60	2.41	13.36	3.22	11.76	5.36	3.85	3.29
11 00.	.00 10	.00						.00	.00	.00		.00	.00			.00	.00 1	.00 1								.00	00					.00		.00 12			.00	.00 12		.00
	· ~	-	-	7	-	7	7	1	-	1	7	-	Н		Н	-	Н	-	-	Н	-	٦	_		_	-	_	-	_	1	-	-	7	-	-	-	.i	7	٦	1
46.700	45.455	46.293	46.380	43.991	13.110	41.647	10.907	40.109	40.068	9.363	46.801	47.653	9.051	9.634	17.750	6.379	5.665	6.267	19.542	50.940	50.969	1.783	51.499	52.820	51.736	50.033	49.960	48.964	49.093	49.553	50.584	51.808	50,295	52.724	51.232	52.463	7.976	6.907	7.187	7.612
		•		4									4	4	4	4	4.	4											-		-		-				55 4	4	4	4
68.108	70.24	71.071	70.62	70.770	69.717	70.173	70.34	69.361	68.126	69.593	72.204	73.001	72.3	72.565	74.4	75.238	74.831	76.242	71.5	71.075	69.74	69.52	70.961	70.17	72.364	68.828	67.5	67.464	66.540	66.455	66.334	65.683	66.865	65.579	66.76	66.104	68.355	68.234	69.187	68.676
964									90,	122	172					93	991																			380	95	128	04	147
34.9	41.3	42.3	43.4	41.3	40.6	40.4	41.7	42.2	41.7	43.3	41.9	42.8	42.9	44.0	42.4	42.5	43.4	41.7	42.0	42.1	42.8	43.7	40.7	40.7	40.1	42.4	43.1	44.2	45.0	42.0	40.9	41.1							46.5	47.5
290	291	291	291	291	291	291	291	291	291	291	292	292	292	292	292	292	292	292	293	293	293	293	293	293	293	294	294	294	294	294	294	294	294	294	294	294	295	295	295	295
ILE A	RG A	RG A	RG A	RG A	RG A	RG A	RG A	RG A	RG A		SN A																HE A							HE A	HE A	HE A	LY A	LY A	LY A	LY A
CDI																																							G	o o
																																							47 (	48 (
2307	3 5	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23	23

7	. 6	9	89	9	9	<b>&amp;</b>	7	7	9	9	œ	9	9	9	9	7	9	9	<b>∞</b>	7	2	9	7	9	9	œ	9	9	80	7	7	9	9	<b>∞</b>	9	9	9	<b>~</b>	7	7
1.00 17.24	1.00 12.68	1.00 13.44	1.00 13.40		1.00 20.79	1.00 23.19	1.00 24.17		1.00 11.38			1.00 12.11	1.00 12.14	1.00 12.77	1.00 13.86	1.00 10.38	1.00 10.10		1.00 13.14								1.00 14.00						1.00 13.50	1.00 14.03	1.00 14.99	1.00 21.08	1.00 21.08	1.00 25.61	1.00 25.23	1.00 11.65
58.967	54.848	53.499	53.225	54.774	56.143	57.233	56.155	52.589	51.254	51.272	50.512	50.252	48.767	48.246	48.324	52.152	52.340	52.822	52.321	53.406	53.770	52.727	53.784	54.264	53.176	53.030	55.470	56.684	56.748	57.595	52.426	51.422	50.319	49.942	50.869	49.859	49.819	50.019	49.600	49.792
74.651	77.036	76.830	77.409	78.176	78.686	78.499	79.477	76.105	75.915	74.957	75.156	75.431	75.462	77.075	78.010	73.951	73.136	74.049	73.974	72.052	71.356	71.038	74.952	75.848	76.824	77.017	76.680	75.761	74.725	76.208	77.423	78.422	77.754	78.286	79.123	80.214	81.169	80.762	82.401	76.630
31.910	32.418	31.740	30.672	33.438	33.863	33.275	34.959	32.405	31.750	30.590	29.639	32.849	32.375	31.759	33.282	30.592	29.383	28.272	27.153	29.712	28.388	30.641	28.541	27.479	27.073	25.875	27.997	28.109	27.432	28.967	27.970	27.549	26.734	25.672	28.821	28.507	29.734	30.875		27.238
ASN A 305 ASN A 306	Ø	Ø		Ø	Ø	Ą	ASN A 306	Ø	K	K	MET A 307	4	K	Ø		K	æ	K	VAL A 308	Ø	æ	∢	ASN A 309	æ	Ø	æ		æ	Ø		Ø	ø	GLN A 310	GLN A 310	Ą	ø	GLN A 310	GLN A 310	Ø	THR A 311
ND2	g	ပ	0	ප	පු	001	ND2	z	S	ပ	0	8	9	SD	S	z	ర	ပ	0	ප	g	CG	z	ర్	ပ	0	ප	ဗ	00	ND2	z	ð	ပ	0	CB	ខ	8	OE1	NE2	z
2433	2435	2436	2437	2438	2439	2440	2441	2442	2443	2444	2445	2446	2447	2448	2449	2450	2451	2452	2453	2454	2455	2456	2457	2458	2459	2460	2461	2462	2463	2464	2465	2466	2467	2468	2469	2470	2471	2472	2473	2474
<b>6</b> 6	æ	9	9	16	9	7	9	9	80	9	9	9	9	9	φ	9	80	1	9	9	69	9	9	80	ထ	7	9	9	90	9	9	9	9	7	9	9	8	9	9	œ
1.00 12.22	. ,	1.00 12.80		1.00 12.28	14	1.00 12.00		1.00 11.89	1.00 12.65		1.00 11.51	1.00 16.19	1.00 17.97	1.00 18.06	1.00 15.93		1.00 21.46						14				1.00 11.06					1.00 12.57	1.00 12.54	1.00 11.76	1.00 12.74	1.00 11.54	1.00 13.50	1.00 12.22	1.00 12.84	1.00 13.98
56.650		57.800	57.393	55.895	56.047	57.555	57.759	56.430	56.325	58.505	59.983	60.914	60.344	62.236	61.683	62.567	63.905	55.418	54.104	53.492	52.785	53.133	53.654	53.849	53.832	53.720	53.156	53.841	53.155	53.367	52.633	51.112	52.971	55.180	55.887	55.479	55.259	57.405		58.097
72.123		71.065	69.787	69.014	67.366	74.374	75.453	75.974	76.181	76.626	76.271	76.197	76.057	75.884	75.723	75.670	75.355	76.026	76.434	75.409	75.782	76.666	77.734	78.877	77,435	74.152	73.086	73.181	73.111	71.742	70.566	70.797	69.248	73.255	73.422	74.720	74.673	73.448	73.551	72.669
38.977	36.791	38.935	39.707	39.027	39.724	38.408	37.462	36.898	35.694	38.131	38.409	37.375	39.739	37.736	40.062	39.029	39.409	37.763	37.256	36.285	35.330	38.419	39.386	38.879	40.573	36.602	35.713	34.348	33.315	36.393	35.687	35.737	36.397	34.290	32.999	32.308	31.100	33.331	32.014	31.174
CA MET A 301 C MET A 301	MET A		CG MET A 301	MET A	MET A	TYR A	TYR A	TYR A	TYR A	TYR A	TYR A	TYR A	TYR A	TYR A	TYR A	TYR A	TYR A	ASP A	ASP A	ASP A	ASP A	ASP A	ASP A	ASP A	ASP A	LEU A	LEU A	LEU A	LEU A		LEU A	LEU A	LEU A	ASN A	CG ASN A 305	ASN A				
2391							_	_		~	_	4	2	9	7	8	σ.	0	-	2412	m	4	'n	۱n	_	80	6	2420	-4	O.	_	₹#	2425	2426	7	_	2429	2430		2432

<b>69</b>	9	9	9	7	7	9	9	<b>&amp;</b>	9	9	9	9	9	9	9	80	7	9	9	7 ه	3	9	9	9	7	7	9	9	8	9	9	9	89	80	7	9	9	ω	9	9
.00 14.60	.00 18.12			.00 22.20	13	.00 11.39	2								.00 15.19			.00 10.64					.00 19.29	.00 19.13	.00 20.57	.00 9.32				.00 12.74			.00 14.20	.00 14.17	9.38	77.6 00.		.00 10.93	.00 10.57	.00 14.03
47.475 1.45.485 1.	160	_	-	7	Н	15.445 1	-	-	157	004 1	915	030	1118	890 1	Н	~	~	_	7	-	7	49.451 1	-	51.345 1	51.636 1	47.174 1	~	1 (99	504 1	_	-	7	48.818 1	50.839 1	44.664 1	285 1	2.636 1	.442 1	42.402 1	.784 1
72.963 4	990	75.856 4			4.		3	•	•		-	•	•	72.767 4	4,	4	4	4	4	4				-	72.819 5	68.796 4		802 4	627	4	47	353	966	829		67.324 4	67.819 4	. 591	•	. 290
17.175	•	•			•	18.316						15.800					•		22.053						19.066	21.044			21.469						21.033			.736	19.995	.660
LYS A 316 LYS A 316	ø	LYS A 316	æ	ø	Æ	TYR A 317	TYR A 317	TYR A 317	æ	æ		Ø	æ	TYR A 317	<b>TYR A 317</b>	Ø	Æ	Æ		Ø	Ø	Ø		Ø	LYS A 318	GLU A 319	K	Ø		GLU A 319	Ø	Ø	GLU A 319	GLU A 319	ASN A 320	Ø	ASN A 320	ASN A 320	Ø	ASN A 320
o 5	ဗ္ဗ	8	8					0		g				~	CZ										NZ		ర	ပ	0	S	ខ	8	OE1	0E2	z	ర	ပ	0	g	ង
2517	2519	2520	2521	2522	2523	2524	2525	2526	2527	2528	2529	2530	2531	2532	2533	2534	2535	2536	2537	2538	2539	2540	2541	2542	2543	2544	2545	2546	2547	. 2548	2549	2550	2551	2552	2553	2554	2555	2556	2557	2558
9 9	· œ	9	œ	9	7	9	9	œ	7	9	9	œ	ø	9	æ	7	7	ø	9	æ	9	9	9	00	80	7	9	9	æ	9	ø	ø	ø	9	9	9	æ	7	9	9
1.00 11.52	1.00 13.81	1.00 12.52	1.00 13.67	1.00 14.32	1.00 11.69	1.00 14.78	1.00 14.32	1.00 16.33	1.00 13.08	1.00 13.20	1.00 14.89	N	1.00 19.18	1.00 30.91	1.00 30.04	(4)	1.00 12.83		1.00 13.20		1.00 10.17	1.00 12.02	_	1.00 15.22	1.00 24.00	1.00 11.76	1.00 10.85	1.00 12.73	1.00 12.47	1.00 12.39	1.00 10.41	1.00 11.23	1.00 12.52	1.00 12.92	1.00 10.90	1.00 12.76	1.00 13.74	1.00 11.54	1.00 14.39	1.00 15.27
48.711	48.599	48.142		47.375	50.561	51.137	51.223	51.276	51.476	51,535	50.150	50.014	52.204	53.718	54.281	54.360	49.111	47.822	46.889	46.175	47.117	47.842	46.985	46.232	47.284	46.892	45.885	46.506	47.510	45.494	44.699	43.316	45.297	42.586	44.563	43.187	42.517	45.861		46.454
75.983	75.572	74.754	73.909	75.250	75.107	74.642	75.792	75.473	76.998	78.108	78.575	79.097	79.260	79.024	78.310	79.691	78.394	79.060	78.137	78.689	79.394	80.510	80.987	80.206	82.100	76.840	75.979	75.368	48	74.838	94	75.610	75.785	76.166	76.350	76.507	77.056	75.634	75.080	73.555
26.432	24.039	27.143	27.593	28.426	25.096	23.812	22.800	21.573	23.320	22.392	21.980	20.827	23.155	23.210	22.384	24.152	22.802	22.396	21.621	20.733	23.753	24.503	25.715	26.285	26.164	21.992	21.297	20.032	20.140	22.265	23.437	23.270	24.613	24.333	25.686	25.510	26.595	18.895	17.638	17.578
THR A		THR A	THR A	THR A	GLY A	GLY A	GLY A	GLY A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	GLU A	GLU A	GLU A	GLU A	GLU A	GLU A	GLU A	GLU A	GLU A	TYR A	LYS A	LYS A	LYS A											
Q (																																							5 S	
2475	2477	2478	2479	248C	2481	2482	2483	2484	2485	2486	2487	2488	2485	2490	2491	2492	2493	2494	2495	2496	2497	2498	2499	2500	2501	2502	2503	2504	2505	2506	2507	2508	2505	2510	2511	2512	2513	251	251	2516





ø	9	9	s	9	ø	7	ø	ø	80	9	ø	<b>a</b> 0	89	7	φ	ø	80	9	9	80	7	14	9	9	<b>8</b> 0	9	ø	7	9	9	7	7	9	9	æ	9	9	æ	æ	7	9
.60 12.60	.60 10.36		.40 10.36	.40 10.82	.40 13.69	.00 9.63	1.00 10.15	1.00 11.71	1.00 10.86	1.00 12.00	.00 12.49	1.00 12.13	1.00 11.72	1.00 9.05	.00 10.52	96.6 00.	61.11 00.1	.00 11.56	1.00 12.65	1.00 11.48	1.00 8.99	1.00 9.23	.00 11.14	1.00 11.62	00 11.55	.00 10.61	1.00 9.76	1.00 11.58	1.00 11.35	00 12.44	.00 11.88	.00 8.48	.00 9.03	.00 12.15	.00 12.94	.00 11.03	.00 11.95	.00 11.39	.00 11.11	.00 11.19	9:36
44.695 0	43.585 0	43.330 0	44.568 0	43.640 0	44.847 0	40.350 1	39.107 1	37.964 1	38.209 1	39.050	39.089	38.870 1	39.354 1	• •	35.569 1	34.352 1	_	35.432 1					31.958 1		30.450 1	30.844 1		31.572	29.370 1		29.733 1	32.263 1	~	32.960 1	32.883 1	31.837 1	30.620 1	29.617 1	30.712 1	33.857 1	34.857 1
64.325	62.693	64.627	64.127	64.862	62.651	65.680	65.987	65.625	65.102	65.323	63.807	63.207	63.200	65.749	65.359	65.556	65.686	66.162				65.385	65.393	66.119	66.836	64.740	65.577	65.938	66.093	66.668	66.756	67.808	69.155	69.793	71.012	70.091	69.815	69.188	70.243	68.936	69.433
37.288	39.550	38.139	37.386	39.604	37.571	38.028	38.762	37.813	36.678	40.149	40.166	39.080	41.228	38.279	37.486	38.334	39.573	36.177	36.351	37.355	35.314	37.802	38.599	39.037	39.744	37.726	36.511	35.652	36.012	34.648	34.853	38.746	39.181	40.073	40.388	37.955	37.069	37.477	35.891	40.568	41.533
CG2AILE A 325	-	∢	CGIBILE A 325	CG2BILE A 325	CDIBILE A 325	N ASP A 326	CA ASP A 326	C ASP A 326	O ASP A 326	CB ASP A 326	CG ASP A 326	OD1 ASP A 326	OD2 ASP A 326	N ASN A 327	CA ASN A 327	C ASN A 327	ø	ASN A	CG ASN A 327	OD1 ASN A 327	ND2 ASN A 327		A HIS A	C HIS A 328	O HIS A 328	HIS A	HIS A	ND1 HIS A 328	CD2 HIS A 328	HIS A	Ø	N ASP A 329	CA ASP A 329	C ASP A 329	ASP A	CB ASP A 329	CG ASP A 329	OD1 ASP A 329	a	4	CA MET A 330
2601	2602	2599	2600	2601	2602	2603	2604	2605	2606	2607	2608	2609	2610	2611	2612	2613	2614	2615	2616	2617	2618	2619	2620	2621	2622	2623	2624	2625	2626	2627	2628	2629	2630	2631	2632	2633	2634	2635	2636	2637	2638
ω	7	7	9	ø	œ	v	ø	9	9	7	9	9	æ	9	9	v	9	7	9	ø	æ	ø	<b>c</b> c	y	7	ø	ø	œ	9	9	9	છ	9	9	ø	7	ø	9	æ	ø	9
1.00 13.17			1.00 10.60	1.00 10.70	1.00 11.96	1.00 9.53	1.00 9.68	1.00 13.05	1.00 9.85	1.00 9.04	1.00 9.68	1.00 11.13	1.00 10.40	1.00 10.10	1.00 9.96	1.00 12.80	1.00 9.05	1.00 9.87	1.00 8.82	1.00 9.73	1.00 10.64			1.00 10.06	1.00 9.44			1.00 10.73	1.00 9.57			1.00 11.07	1.00 11.10	1.00 11.33	1.00 13.68		1.00 9.24	1.00 9.67			0.60 10.90
43.445	42.323	43.422	42.896	42.685	43.452	44.024	44.124	45.415	42.930	41.566	41.265	41.584	40.935	39.767	39.384	39.396	40.258	42.594	43.130	42.453	41.902	44.652	44.887	45.318	42.559	41.880	42.385	42.805	40.286	39.817	39.810	39.354	39.380	38.926	38.931	42.285	42.710	41.480	41.645	43.596	42.764
	67			67.643	66.685		71.176	71.820	72.064	67.671	66.478	66.821	67.716	66.118	65.842	64.954	64.698	66.133	66.524	65.704	64.628	66.302				65.582	66.205	67.375	65.698	67.122	67.817	67.746	69.142	69.083	69.788	65.385	65.879	66.277	67.116	64.869	63.705
18.619	17.558	23.285	24.610	25.415	25.448	25.299	24.761	25.310	25.183	26.185	27.014	28.477	29.071	26.872	25.387	27.793	24.773	29.019	30.333	31.433	31.218	30.364	30.009	29.314	32,660	33.784	35.086	35.086	33.716	33.638	32.421	34.798	32.321	34.683	33.498	36.137	37.469	38.277	39.255	38.211	38.779
OD1 ASN A 320	K	LEU A	A LEU A	C LEU A 321	O LEU A 321	CB LEU A 321	CG LEU A 321	CD1 LEU A 321	CD2 LEU A 321	ILE A	CA ILE A 322	C ILE A 322	O ILE A 322	CB ILE A 322	CG1 1LE A 322	CG2 ILE A 322	CD1 1LE A 322	N THR A 323	CA THR A 323	THR A	O THR A 323	CB THR A 323	THR A	CG2 THR A 323	N PHE A 324		PHE A	PHE A	CB PHE A 324	PHE A	PHE A		CE1 PHE A 324	CE2 PHE A 324	CZ PHE A 324	ILE A		C ILE A 325	ILE A	CB AILE A 325	
2559	2560	2561	2562	2563	2564	2565	2566	2567	2568	2569	2570	2571	2572	2573	2574	2575	2576	2577	2578	2579	2580	2581	2582	2583	2584	2585	2586	2587	2588	2589	2590	2591	2592	2593	2594	2595	2596	2597	2598	2599	2600



2681
8 2682 CA SER A
6 2683
0 #897 91 15 7876 71
8
7 2687
6 2688
2689
8 2690
6 2691
8 2692
2693
6 2694
6 2695
8 2696
6 2697
6 2698
6 5699
7 2700
6 2701
10.61 7 2702
7 2703
2704
ဖ
9
œ
9
<b>9</b>
9
y Q
9
9
9
7 2715
6 2716
6 2717
8 2718
6 2719
6 2720
6 2721
13.51 6 2722 C

9.98 111.63 111.63 113.09 113.09 113.09 110.65 110.65 110.88 110.25 110.25 110.26 110. 51.003 51.273 51.534 52.310 51.160 51.330 50.717 50.717 51.357 51.357 51.357 52.605 52.605 52.605 52.605 52.605 48.987 48.462 48.339 47.829 48.079 44.722 51.660 59.197 59.764 59.764 59.747 60.725 60.600 61.920 63.143 63.422 64.348 65.659 66.030 66.526 67.747 67.747 68.189 69.115 69.115 69.115 69.115 69.115 62.418 63.250 62.379 62.887 63.827 61.107 60.261 59.999 60.087 58.885 57.574 39.590 33.764 33.764 34.473 35.809 36.473 36.473 36.473 36.809 37.367 38.120 38 27,65 27,66 27,67 27,67 27,77 27 18.65 22.63 112.85 14.45 113.48 113.48 110.37 110.38 110.62 112.01 113.17 113.1 13.40 12.11 9.09 10.04 12.75 11.63 12.61 10.53 10.43 10.42 10.38 11.01 9.73 10.78 12.69 1.00 .039 446.229 445.299 44.550 44.283 48.898 49.570 50.370 50.370 50.439 48.370 44.699 44.564 50.345 51.433 51.233 52.806 50.731 50.731 50.731 50.731 51.233 52.806 52.806 52.806 53 157 024 831 008 324 48. 8 54.662 60.100 61.036 61.899 62.067 54.658 55.099 58.363 58.557 59.830 59.830 60.911 60.911 61.737 64.044 63.859 61.049 60.715 59.935 60.288 60.288 59.854 60.288 58.294 58.977 58.185 59.122 58.916 064 436 580 612 40.189 48.951 42.520 43.612.520 43.612.520 44.114 44.114 40.049 40.049 41.847 43.132 43.232 48.937 50.377 46.585 45.626 44.835 43.661 46.346 

10.12 10.09 10.66 11.54 36.636 37.510 36.796 35.363 40.550 42.925 38.176 36.807 37.653 39.927 37.647 40.671 34.206 32.960 34.066 32.937 31.670 35.418 35.210 35.593 35.297 61.021 59.991 60.760 62.051 61.558 62.183 62.183 62.183 62.183 62.183 62.183 62.183 61.272 61.272 61.273 61.273 61.274 61.274 61.274 61.274 61.274 61.274 61.274 61.274 61.274 62.148 57.488 56.287 56.846 58.150 56.052 58.715 56.601 35.810 36.364 37.027 37.029 37.399 37.193 37.193 39.277 39.123 39.123 39.274 39.123 39.556 39.556 37.937 38.086 37.937 29.245 27.914 30.508 31.003 32.509 33.32.509 33.32.503 34.703 34.703 34.703 34.313 34.313 34.313 34.313 33.858 33.323 38.913 40.155 39.988 2863 2863 2864 2865 2865 2866 2867 2873 2873 2873 2874 2875 2877 2878 2879 2880 2881 2882 2883 2884 2885 2886 2887 2888 2889 2889 2889 2850 2851 2853 2853 2854 2855 2856 2857 2859 2859 2860 2860 10.64 12.51 13.68 13.74 11.34 11.34 11.08 9.52 9.52 10.03 10.73 10.73 11 10.84 11.59 10.71 11.61 12.49 12.69 12.54 12.15 11.14 11.00 10.49 13.18 15.74 10.24 10.01 10.67 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 50.754 50.024 48.533 48.085 50.545 46.595 50.850 51.960 52.149 53.022 53.296 53.420 53.324 51.368 51.432 50.536 49.726 51.012 52.045 50.754 50.024 548 48.317 47.894 47.780 46.318 45.929 688 43.066 43.698 581 769 50.229 49. 42. 70.157 69.113 71.370 60.657 62.022 62.214 60.375 65.403 65.402 65.741 65.920 65.997 66.369 67.185 68.504 68.853 70.130 65.584 64.196 63.299 64.007 62.722 63.621 62.301 62.052 60.948 58.939 64.357 64.574 63.688 65.707 62.648 27.455 27.638 27.867 24.542 24.624 32.492 33.679 31.500 30.267 22.858 30.092 31.244 31.244 31.244 31.244 31.244 32.092 22.100 22.29 30.607 22.29 30.607 22.29 30.607 22.29 30.607 22.29 30.607 22.29 30.607 22.29 30.607 22.29 30.607 22.29 30.607 22.29 30.607 22.29 30.607 22.29 30.607 22.29 30.607 22.29 30.607 22.29 30.607 22.209 22.607 22. 24.669 25.099 26.535 26.964 25.149 25.546 25.885 27.279 27.512 24.908 23.622 

ဖ	9	9	16 6	7	9	• •	oνα	7	9	9	80	٠,		80	7	9	7 و و	8 4	<b>.</b> .	89	7	9	vo c	2 C	7	9	9	80	9	9	89	æ	7	9 '	9
1.00 14.58	11.3	11	1.00 13.40	12.7		1.00 14.06	15.4		1.00 15.04			1.00 12.36		~				1.00 14.23	0.60 25.40	0.60 11.28		0.40 16.79		0.40 23.38	12	13		13	2		12	12	1.00 11.77	1.00 11.70	1.00 11.76
37.586	36.115	35.397	34.139	36.904	37.152	35.754	37.904	35.729	34.408	34.571	35.433	33.872	33.102	32.888	32.710	32.081	30.645	30.084	32.279	31.899	32.756	31.871	32.513	33.793	30.152	.28.792	27.745	27.614	28.451	27.084	26.464	26.659	27.151	26.928	27.918
58.365	57.163	55.810	56.012 56.859	58.817	60.193	60.849	60.184	62.152	62.780	64.242	64.643	65.045	66.898	66.015	68.167	68.579	68.064	67.570	70.451	69.778	71.695	70.115	70.670	70.909	68.230	67.780	67.942	69.093	68.624	68.351	67.384	69.190	66.874	65.582	64.544
46.195	44.602	44.316	42.994	47.271	47.689	47.745	49.047	47.908	48.101	47.715	46.895	48.524	47.034	46.202	46.994	45.708	45.594	46.556	43.956	43.002	43.728	45.872	44.590	43.560	44.373	44.018	45.054	45.503	42.737	42.153	42.556	41.293	45.561	44.932	45.436
MET A 367 MET A 367	Ø	A 3	SD MET A 367 CE MET A 367	Ø	4	<b>A</b> A		ø	æ	GLY A 369	Æ	æ ,	GLY A 370	4	æ	<b>4</b>	K ,	ASN A 371	. ∢	A	ø		<b>A</b>	UDIBASN A 371 ND2BASN A 371		ASP A	ASP A	ASP A	ASP A	ASP A	1 ASP A 372		PRO A		PRO A
2933 C 2934 O			2937 SD 2938 CE										2950 C						2957 CG			2956 CB			2960 N						2966 OD1	2967 002	_	2969 CA	2970 C
29	29	29	29	29	29	5 6	2 6	29	29	29	29	23	2 2	29	29	53	29	23	2 62	29	29	29	5 5	2 62	29	29	29	29	29	29	29	29	29	5 5	29
99	ω	7	φφ			oo u	۰ ۲	9	9	60	9	φ (	0 00	ο ο	7	9		ω v	o vo	9	œ	7	7	o o	80	9	9		_	ဖ		9	89	7	9
0.04	36	S	.33	. 29	07	12.04	11.97	11.29	12.77	14.39	10.27	1.76	13.43	11.75	0.80	11.28	2.82	12.70	9.91	14.70	17.54	12.05	10.84	13.18	14.90	1.57	1.25	14.66	3.34	15.95	14.89	12.93	12.26	12.15	10.89
1.00 12	12.	12	1.00 11.				1.00						1.00.1					. 00		1.00				1.00 1					1.00 13	1.00 15	1.00 1	1.00 1	1.00	1.00	1.00
	1.00 12.	1.00 12	= =	1.00	7 1.00		1.00	1.00		1.00	1.00	1.00		1.00	1.00	1.00	7.00	7.00		1.00	1.00		1.00	1.00	1.00	1.00	1.00	1.00		1.00			46.302 1.00	38.357 1.00	37.181 1.00
513 1.00	38.093 1.00 12.	38.446 1.00 12	39.736 1.00 11 40.693 1.00 11	41.376 1.00	40.387 1.00	39.494 1.00	40.670 1.00	41.610 1.00	41.198 1.00	41.915 1.00	41.712 1.00	40.469 1.00	39.239 1.00	38.247 1.00	39.993 1.00	39.542 1.00	39.629 1.00	39.219 1.00	1.00	38.693 1.00	1.00	38.847 1.00	40.333 1.00	1.00	39.205 1.00	41.514 1.00	326 42.813 1.00	215 43.707 1.00	977 43.139 1.00	747 44.882 1.00	.332 1.00	395 45.154 1.00	52.939 46.302 1.00	.315 38.357 1.00	56.998 37.181 1.00
55.099 36.513 1.00 55.744 37.772 1.00	8 55.431 38.093 1.00 12.	2 56.614 38.446 1.00 12	6 57.169 39.736 1.00 11 3 56.009 40.693 1.00 11	2 56.074 41.376 1.00	8 57.957 40.387 1.00	8 59.065 39.494 1.00	1 54.961 40.670 1.00	9 53.835 41.610 1.00	0 52.938 41.198 1.00	5 51.968 41.915 1.00	14 53.093 41.712 1.00	52.265 40.469 1.00	19 53:001 39:248 1:00 12 54:257 39:239 3:00	3 52.343 38.247 1.00	6 53.105 39.993 1.00	6 52.398 39.542 1.00	32 53.305 39.629 1.00	16 52.953 39.219 1.00	4 50.951 37.828 1.00	13 49.690 38.693 1.00	6 49.200 39.269 1.00	4 49.142 38.847 1.00	7 54.426 40.333 1.00	13 55.357 40.582 1.00	18 56.182 39.205 1.00	7 54.753 41.514 1.00	72 54.326 42.813 1.00	0 55.215 43.707 1.00	2 52.977 43.139 1.00	3 54.747 44.882 1.00	1 52.494 44.332 1.00	1 53.395 45.154 1.00	18 52.939 4	56.315 38.357 1.00	00 56.998
GLY A 362 40.454 55.099 36.513 1.00 GLY A 362 40.997 55.744 37.772 1.00	GLY A 362 42.168 55.431 38.093 1.00 12.	THR A 363 40.222 56.614 38.446 1.00 12	THR A 363 40.676 57.169 39.736 1.00 11 THR A 363 41.033 56.009 40.693 1.00 11	THR A 363 42.072 56.074 41.376 1.00	THR A 363 39.528 57.957 40.387 1.00	THR A 363 39.248 59.065 39.494 1.00	THK A 363 40.024 58.554 41.730 1.00 GLU A 364 40.221 54.961 40.670 1.00	GLU A 364 40.379 53.835 41.610 1.00	GLU A 364 41.520 52.938 41.198 1.00	GLU A 364 41.845 51.968 41.915 1.00	GLU A 364 38.994 53.093 41.712 1.00	GLU A 364 38.652 52.265 40.469 1.00	GLU A 364 38.139 53.001 39.246 1.00	GLU A 364 37.813 52.343 38.247 1.00	GLN A 365 42.046 53.105 39.993 1.00	GLN A 365 43.256 52.398 39.542 1.00	GLN A 365 44.462 53.305 39.629 1.00	GLN A 365 45.606 52.953 39.219 1.00	GLN A 365 43.136 51.923 30.000 1.00 GLN A 365 41.964 50.951 37.828 1.00	GLN A 365 42.043 49.690 38.693 1.00	GLN A 365 41.016 49.200 39.269 1.00	GLN A 365 43.204 49.142 38.847 1.00	TYR A 366 44.317 54.426 40.333 1.00	TYR A 356 45.443 55.357 40.582 1.00 TYR A 366 46.039 55.921 39.308 1.00	TYR A 366 47.248 56.182 39.205 1.00	TYR A 366 46.547 54.753 41.514 1.00	TYR A 366 45.872 54.326 42.813 1.00	TYR A 366 45.270 55.215 43.707 1.00	TYR A 366 45.942 52.977 43.139 1.00	TYR A 366 44.653 54.747 44.882 1.00	TYR A 366 45.361 52.494 44.332 1.00	TYR A 366 44.711 53.395 45.154 1.00	TYR A 366 44.118 52.939 4	MET A 367 45.147 56.315 38.357 1.00	MET A 367 45.700 56.998
GLY A 362 40.454 55.099 36.513 1.00 GLY A 362 40.997 55.744 37.772 1.00	O GLY A 362 42.168 55.431 38.093 1.00 12.	N THR A 363 40.222 56.614 38.446 1.00 12	A 363 40.676 57.169 39.736 1.00 11 A 363 41.033 56.009 40.693 1.00 11	O THR A 363 42.072 56.074 41.376 1.00	CB THR A 363 39.528 57.957 40.387 1.00	OG1 THR A 363 39.248 59.065 39.494 1.00	THK A 363 40.024 58.554 41.730 1.00 GLU A 364 40.221 54.961 40.670 1.00	CA GLU A 364 40.379 53.835 41.610 1.00	C GLU A 364 41.520 52.938 41.198 1.00	O GLU A 364 41.845 51.968 41.915 1.00	CB GLU A 364 38.994 53.093 41.712 1.00	CG GLU A 364 38.652 52.265 40.469 1.00	CD GDU A 364 38.139 53.001 35.246 1.00	OE2 GLU A 364 37.813 52.343 38.247 1.00	N GLN A 365 42.046 53.105 39.993 1.00	CA GLN A 365 43.256 52.398 39.542 1.00	C GLN A 365 44.462 53.305 39.629 1.00	O GLN A 365 45.606 52.953 39.219 1.00	CG GLN A 365 43.139 51.523 58.009 1.00	CD GLN A 365 42.043 49.690 38.693 1.00	OE1 GLN A 365 41.016 49.200 39.269 1.00	NE2 GLN A 365 43.204 49.142 38.847 1.00	N TYR A 366 44.317 54.426 40.333 1.00	TYR A 356 45.443 55.357 40.582 1.00 TYR A 366 46.039 55.921 39.308 1.00	O TYR A 366 47.248 56.182 39.205 1.00	CB TYR A 366 46.547 54.753 41.514 1.00	CG TYR A 366 45.872 54.326 42.813 1.00	CD1 TYR A 366 45.270 55.215 43.707 1.00	CD2 TYR A 366 45.942 52.977 43.139 1.00	CE1 TYR A 366 44.653 54.747 44.882 1.00	A 366 45.361 52.494 44.332 1.00	CZ TYR A 366 44.711 53.395 45.154 1.00	OH TYR A 366 44.118 52.939 4	MET A 367 45.147 56.315 38.357 1.00	CA MET A 367 45.700 56.998

9 4 9 9 9 9 9 13.09 13.28 13.26 12.95 14.01 17.99 17.24 14.68 16.71 14.00 15.78 13.86 13.98 13.69 17.55 57 34.636 33.751 33.975 34.625 35.190 30.812 31.558 32.790 34.277 35.254 32.750 32.726 33.204 32.914 35.761 36.094 669. 30.804 27.749 27.113 25.713 25.530 30.324 31.637 31.705 669 35.052 31.670 37.641 826 33.775 31.613 33. 53.836 55.572 57.572 57.572 57.572 57.573 57.572 57.573 57.672 46.466 45.806 46.669 46.629 47.618 47.480 48.469 48.395 46.138 .402 .732 .868 .826 44.714
44.714
41.714
41.543.750
42.817
41.543.364
42.364
43.364
40.896
40.855
40.855
40.856
40.856
40.856
40.856
40.856
40.856
40.856
40.856
40.856
40.856 44.854 42.029 40.701 40.791 39.963 39.742 39.619 572 327 38.414 39.293 41.721 42.065 41.067 40. 3039 3041 3042 3043 3044 1045 3046 047 11.30 10.91 13.69 9.68 9.68 11.01 11.01 12.69 14.25 14.25 14.25 19.98 19.98 19.98 19.01 10.01 10.01 10.01 10.01 10.01 10.01 10.01 10.01 10.01 10.01 10.01 10.01 10.01 10.01 10.01 10 12.18 10.84 13.47 14.66 11.13 11.23 12.81 111.47 111.36 111.93 111.44 113.66 113.05 113.05 113.05 10.05 10.05 10.05 110.0 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 25.479 25.475 26.040 28.815 30.741 31.346 30.101 28.884 27.954 27.954 27.956 26.613 31.004 31.991 31.991 31.991 31.991 32.073 32.073 33.073 33.073 33.073 34.073 35.073 37 .749 .407 .297 .658 825 646 638 25. 66.349 63.895 63.320 62.353 64.456 64.456 65.356 65.147 66.403 62.009 65.136 65.136 65.755 64.933 64.933 66.338 66.051 66.051 66.338 66.051 66.338 66.051 66 59.886 61.789 62.997 63.980 65.214 66.011 67.071 59.361 58.173 56.943 56.949 51.271 45.203 44.295 43.275 42.406 43.489 42.823 42.889 42.188 43.519 43.410 40.908 40.177 39.567 39.569 45.014 46.632 46.484 46.484 47.139 447.139 46.896 48.493 49.258 49.457 50.385 50.125 50.572 41.975 .944 44.270 

<b>9</b> 9	99	7	9	9	<b>&amp;</b>	9	9	9	9	7	7	9	9	8	9	9	9	œ	8	70	9	9	80	9	9	9	7	9	9	ထ	9	80	7	9	9	ထ	9	80	9
1.00 14.39	1.00 12.44	12.	12		1.00 12.66	12.	15.					1.00 11.43				1.00 10.67			12.						1.00 13.75				1.00 10.96	1.00 12.48		1.00 19.93	1.00 11.04	1.00 12.34		1.00 11.63	1.00 12.40	1.00 14.67	1.00 14.84
41.777	41.216	46.775	ഹ	48.699	49.197	48.917	48.243	49.325	49.440	50.107	48.782	49.475	48.742	49.337	49.487	49.997	49.810	49.226	50.284	47.403	46.653	46.716	46.930	45.175	44.296	45.061	46.628	46.686	48.021	48.186	46.394	46.423	49.099	50.445	50.682	51.169	51.509	51.401	52.927
48.299	47.536		•	•	49.556	48.000	46.712	45.970	46.524	45.465	50.908	52.048	52.482	52.735	53.223	54.519	55.744	55.628	56.841	52.675	53.176	52.128	52.481	53.410	53.661	54.570	50.846	49.822	49.782	49.774	48.431	47.538	49.942	49.922	51.130	51.097	49.889	48.614	50.011
37.375	36.340	38.053	37.303	36.544	35.438	38.291	38.798	39.589	40.980	41.835	37.256	36.636	35.375	34.300	37,670	36.957	37.870	38.974	37.458	35.455	34.279	33.156	31.968	34.667	33.434	35.702	33.487	32.426	31.708	30.469	33.059	31.944	32.493	31.920	31.061	29.935	33.039	33.699	32.443
CD2 PHE A 389 CE1 PHE A 389	CE2 PHE A 389	LYS A	LYS A	æ	LYS A	LYS A	LYS A	LYS A	LYS A	LYS	ø	GLU A	GLU A		GLU A	4	GLU A	GLU A	GIN A		VAL A	ď	VAL A	VAL A	VAL A	32 VAL A	SER A	A SER A	æ	SER A	SER A		THR A	CA THR A 394		O THR A 394	THR A	-	CG2 THR A 394
3097 3098	3099	3101	3102	3103	3104	3105	3106	3107	3108	3109	3110	3111	3112	3113	3114	3115	3116	3117	3118	3119	3120	3121	3122	3123	3124	3125	3126	3127	3128	3129	3130	3131	3132	3133	3134	3135	3136	3137	3138
ω α	۷ ع	· v	œ	ø	œ	9	7	9	9	œ	9	80	9	7	9	9	80	9	80	9	7	9	9	æ	9	8	9	7	9	9	80	9	7	9	9	œ	9	9	9
1.00 20.99	1.00 13.75	13.7	18.1			20.2		1.00 16.71	1.00 16.63	1.00 14.22	1.00 19.15	1.00 17.99	1.00 24.33	1.00 14.92	1.00 13.63	1.00 11.22	12			11	1.00 14.21	11		1.00 12.57	1.00 17.16		14		10	1.00 12.23	Ξ	1.00 10.26	1.00 12.53	1.00 13.64	1.00 13.47	1.00 12.37		1.00 12.01	1.00 16.87
41.805	40.950	43.345	44.298		41.937	40.342	43.573	44.906	45.454	46.539	45.009	44.288	44.541	44.606	45.131	46.328	46.478	44.054	43.822	42.719	47.184	48.316	47.887	48.437	49.145	49.669	50.273	46.838	46.360	45.808	46.001	45.239	45.084	44.579	45.779	45.789	43.763	43.141	43.916
44.428		44	44.	43.	42.	43.			45.974	45.952	43.546	43.878	42.234	47.039	48.254	48.801	48.649	49.344	49.919	48.809	49.552	50.167	51.134	51.113	50.926	49.939	51.724	51.910	52.785	52.083	52.497	53.700	50.956	50.182	49.746	49.895	48.975	48.178	47.248
43.286	40.302	39.773	38.951	38.098	38.725	37.515	41.088	41.648	42.313	42.873	42.693	43.883	42.075	42.254	42.952	42.175	40.990	43.101	41.805	43.656	42.886	42.237	41.127	40.045	43.277	44.177	42.644	41.441	40.376	39.162	38.030	40.968	39.347	38.202	37.361	36.157	38.766	37.627	36.936
ASP	A 4	A	K	THR A	THR A	THR A	THR A	THR A	THR A	THR A	THR A	THR A	THR A	THR A	THR A	THR A	THR A	THR A	THR A	4	THR A	THR A	THR A	THR A	THR A	THR A	THR A	ALA A	Ø	ALA A	ALA A	ALA A	PHE A	PHE A	PHE A	PHE A	PHE A	PHE A	PHE A
007	z ć	່ວ	0	CB	061	g	z		3066 C																	3083 0											3094 CE		

,81 ,9 16.08 14.78 14.12 115.84 14.12 14.12 14.12 14.12 17.91 11.03 11.03 11.48 11.48 11.48 11.98 53.373 54.506 53.980 55.3980 55.311 56.564 56.564 57.762 54.104 57.762 5 46.399 46.343 45.640 46.978 50.972 52.060 53.189 52.590 51.404 50.797 51.102 50.634 49.2179 51.128 49.2179 52.138 52.358 52.828 52.828 52.828 52.828 63.948 64.9.617 64.9.617 64.9.617 65.93 66.93 6 58.973 58.253 59.410 59.266 60.152 60.545 23.412 25.452 25.452 26.214 27.124 27.127 27.207 27 CAN NO CA 31881 31882 31883 31884 31886 12.76 11.35 10.18 11.69 11.69 11.59 11.70 11.09 11.42 10.92 14.21 10.96 11.76 12.24 12.24 12.96 13.71 17.33 14.09 50.842 51.404 52.491 51.900 52.434 53.229 54.353 55.479 56.754 50.754 557 163 075 203 825 825 510 656 49.068 48.158 182 688 446 436 322 592 624 624 976 604 811 4 54.747 53.116 53.116 53.118 52.318 52.318 52.318 52.32 51.24 52.230 52.230 52.231 53.871 52.833 53.871 54.478 54.478 55.603 56.271 57.2788 57.278 57.278 57.278 57.278 57.278 57.278 57.278 57.278 57.2788 57.278 57.278 57.278 57.278 57.278 57.278 57.278 57.278 57.2788 57.278 57.278 57.278 57.278 57.278 57.278 57.278 57.278 57.2788 57.278 57.278 57.278 57.278 57.278 57.278 57.278 57.278 57.2788 57.278 57.278 57.278 57.278 57.278 57.278 57.278 57.278 57.2788 57.278 57.278 57.278 57.278 57.278 57.278 57.278 57.278 57.2788 57.278 57.2788 57.27 52.334 53.558 53.709 54.373 54.804 54.745 31.600 32.9.666 33.0.784 33.0.784 33.0.676 33.0. 

82 15.64 14.70 222.99 222.07 119.41 12.19.41 12.19.41 11.10.19 13.09 13.13.13 14.79 19.43 110.13 14.89 15. 15. 64.700 66.510 64.562 65.579 580 67.503 66.470 64.163 63.633 63.330 65.082 63.947 62.833 65.651 66.709 67.898 62.991 61.110 61.592 63.062 63.824 61.335 64.474 68.075 66.591 68.773 53.038 65.507 55. 69.676
70.302
70.153
70.153
71.623
71.6123
71.6123
71.6123
71.6123
68.430
68.482
66.786
66.983
68.784
66.786
66.983
68.785
69.784
66.786
67.884
66.7884
66.7884
66.883
68.782
69.888 71.491 72.134 71.827 72.197 73.061 36.215 37.012 38.296 38.300 25.620 29.899 29.899 31.959 31.959 31.959 31.959 31.959 31.959 32.409 32.409 33.759 34.459 35.259 35.259 37.112 36.113 37.112 37.113 37.113 37.113 37.113 37.113 37.113 37.113 37.113 37.113 37.113 37.113 37.113 37.113 37.113 310 44112 44112 44112 44112 44112 44112 44113 44113 44114 4414 32665 32665 32667 32667 32707 32 11.82 12.00 12.00 12.00 13.00 13.00 13.00 14.00 11.00 1.00 1.00 1.00 1.00 1.00 1.00 49.360 49.869 49.549 49.395 49.395 51.499 51.702 52.068 51.144 49.759 51.088 51.144 49.515 48.707 53.678 55.005 55.030 54.133 56.207 56.558 56.707 56.777 57.928 57.928 57.762 56.790 57.141 48.918 50.894 58.225 54.786 56.192 .919 58.321 342 952 55 61.748 60.418 60.742 59.420 59.595 58.598 63.743 64.361 65.790 66.253 66.393 67.645 67.370 68.210 67.169 68.677 68.449 68.261 68.740 67.959 68.392 68.665 69.430 69.192 59.494 58.136 56.983 55.775 57.356 62.154 64.182 65.190 62.678 61.567 69.229 70.423 66.229 891 20.265 19.875 20.511 19.521 19.521 18.272 19.757 18.848 19.168 117.440 116.935 116.935 116.937 116.931 116.441 116.441 116.441 116.433 20.156 20.277 20.277 20.273 21.566 21.566 21.362 22.822 23.822 20.812 27.425 301 

83 15.49 11.61 19.26 11.61 10.90 11.03 11.03 11.03 11.05 57.021 556.683 61.025 61.458 61.282 61.282 62.193 63.133 63.133 63.133 63.133 63.133 63.133 63.133 64.484 60.25 60.004 60.276 60.2776 60.277 58.643 .620 .805 699 847 58 69.780 70.412 66.508 67.148 67.148 67.148 68.198 68.198 68.198 68.198 69.198 32.692 31.503 31.503 32.649 32.649 32.649 33.251 33 33349 33351 333521 333521 33352 33352 33362 33362 33362 33362 33362 33370 33370 33372 33374 3379 3380 3381 3382 3383 3384 3386 3386 18.18 113.47 114.63 115.76 116.07 112.99 113.34 113.34 115.09 117.99 113.34 119.99 119.99 110.52 110.52 110.52 110.52 110.52 111.52 111.52 111.52 111.52 111.52 111.52 111.53 111.53 111.53 111.54 111.55 111 1.00 62.444 59.764 59.636 59.977 59.383 58.138 57.856 58.754 59.854 62.797 61.526 60.536 60.536 63.699 62.591 64.597 61.313 59.907 59.907 59.908 59.909 59.909 59.808 59.808 59.808 59.808 60.338 60.338 60.348 60.348 62.145 62.481 419.846 90 72.686 71.266 71.266 71.152 71.879 71.879 71.879 71.979 69.947 72.836 71.526 71 66.958 65.959 67.958 66.266 67.172 67.548 66.328 65.254 67.866 35.862 39.349 40.517 41.103 40.399 42.250 42.250 42.129 42.755 42.757 42.757 44.190 43.170 43.170 43.109 44.1295 42.126 44.120 44.520 44.520 46.759 41.304 40.281 38.910 38.910 39.231 41.737 36.628 35.772 35.908 35.908 36.410 35.063 34.258  $\begin{array}{c} 0.00\\$ 

84 14.21 14.56 14.56 11.58 11.58 11.158 11.153 12.54 10.81 10.81 10.81 11.254 11.255 11.254 11.255 11.2 11.73 9.07 9.49 10.58 60.800 61.285 60.583 60.577 59.912 59.603 57.905 57.615 60.522 62.427 62.734 63.033 60.583 60.577 62.831 63.562 510 044 59.131 59.702 59.844 59.967 58.471 58.255 58.940 57.490 60.600 60.876 59.826 59.978 60.409 59.363 57.400 674 812 072 017 63. 65. 57.501 56.561 56.561 56.561 56.561 56.7501 56.7501 57.602 58.750 59.750 59.750 59.750 59.750 59.750 59.750 59.750 60.140 20.026 18.927 117.727 16.804 22.538 22.538 23.993 22.632 23.993 22.632 23.993 24.721 26.043 27.057 26.043 27.057 26.043 27.057 2 3465 3466 3467 3468 3433 3435 3435 3435 3436 3439 3440 3441 3442 3442 3443 3444 11. 90 11. 50 11. 50 11. 53 11. 53 11. 53 11. 56 11. 56 11. 50 11. 55 11. 55 11. 57 11 1.00 11.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 58.542 59.290 60.626 61.643 62.643 63.865 62.639 62.639 63.008 63.733 64.570 63.325 60.547 59.986 62.659 60.300 62.715 60.345 61.552 63.623 63.080 63.233 64.340 63.854 63.271 63.342 62.144 62.238 61.481 60.252 61.450 61.449 62.357 61.773 60.836 61.354 61.132 62.175 62.918 62.396 63.858 63.336 64.087 59.522 58.735 57.587 57.591 58.214 58.370 58.370 57.810 58.173 59.092 22.068 20.937 20.937 20.937 20.931 20 13.018 11.168 12.246 11.311 15.546 16.284 17.573 15.484 14.267 14.452 13.103 

<b>a</b> o vo	9	9	80	7	7	9	9	89	ø	8	7	9	9	80	9	8	7	9	9	8	5	9	9	9	9	9	9	8	7	9	9	80	9	œ	7	9	9	œ	9	9
1.00 18.55		1.00 48.48		1.00 54.59	1.00 13.09	1.00 13.14		1.00 15.27		1.00 17.34	1.00 13.46	1.00 13.64	1.00 14.33	1.00 15.22			1.00 13.24	1.00 12.26			1.00 13.59		1.00 12.95		1.00 11.89	1.00 11.54	1.00 12.16	1.00 15.42	1.00 14.60	1.00 14.67	1.00 13.95	1.00 16.35	1.00 17.27	1.00 18.88	1.00 15.00	1.00 13.10	1.00 14.11	1.00 17.51	1.00 14.59	1.00 15.39
61.812	57.761	57.810	58.444	57.127	61.192	62.559	62.958	62.168	62.702	62.516	64.198	64.730	65.599	65.974	65.791	65.203	65.817	66.694	67.749	67.451	65.860	62.009	65.629	63.667	64.882	62.914	63.551	62.811	69.008	70.040	69.912	69.879	71.418	71.733	69.758	69.559	70.863	71.131	68.403	67.136
62.543	62.971	61.634	61.462	60.605	63.161	63.516	62.739	62.718	65.025	65.770	62.251	61.703	62.835	63.761	60.598	59.420	62.845	63.886	63.156	62.293	64.824	65.807	66.940	65.549	67.843	66.474	67.595	68.508	63.663	63.232	64.002	65.229	63.684	62.764	63.265	63.892	63.757	62.703	63.188	63.120
52.198	53.543	54.248	55.292	53.729	50.115	49.738	48.481	47.524	49.451	50.658	48.417	47.177	46.581	47.366	47.452	48.033	45.302	44.693	43.877	43.032	43.772	44.592	45.124	44.836	45.908	45.595	46.107	46.871	43.976	43.034	41.698	41.739	43.620	44.701	40.628	39.303	38.498	37.935	38.537	39.386
O GLN A 440 CB GLN A 440	_	GLN A	GLN A	NE2 GLN A 440				O SER A 441								G SER A 442		CA TYR A 443	TYR A 443				D1 TYR A 443				CZ TYR A 443								ILE A 445	A ILE A 445		•	ILE A	CG1 ILE A 445
3517 O 3518 C					3523 N	3524 C	3525 C		3527 C		3529 N										3539 C									3548 C						3554 C		3556 0	3557 C	_
<b>ω</b> φ	9	9	9	7	9	9	60	9	9	80	7	7	9	9	œ	9	9	9	7	9	7	7	7	9	9	8	9	ų	80	7	7	9	9	80	9	80	9	7	9	9
1.00 10.88	1.00 11.49	1.00 10.37	1.00 13.59	1.00 10.83	1.00 11.64	1.00 9.83	1.00 12.85	1.00 12.59	1.00 12.87	1.00 13.07	1.00 13.56	1.00 10.03	1.00 9.39	1.00 10.71	1.00 12.32	1.00 12.80	1.00 13.58	1.00 11.13	1.00 10.90		1.00 11.67	1.00 13.84	1.00 11.11	1.00 12.20	1.00 13.73	1.00 12.56	1.00 13.49	1.00 13.92	1.00 15.45	1.00 13.51	1.00 11.49	1.00 12.77	1.00 14.69	1.00 16.23	1.00 16.48	1.00 17.73	1.00 15.47	1.00 14.00	1.00 16.61	1.00 19.80
58.922	56.120	55.689	54.758	58.596	58.943	57.715	57.309	60.036	60.383	60.026	61.224	57.181	55.958	56.267	55.310	55.161	55.869	55.043	55.738	55.195	53.948	55.947	57.513	57.734	57.869	58.830	59.071	59.255	58.985	59.700	56.949	56.883	57.846	57.917	55.456	54.990	54.554	58.501	59.481	60.910
64.734	62.608	63.352	63.228	62.933	63.646	63.653	62,621	62.821	63.480	64.602	62.721	64.891	64.887	64.725	64.552	66.201	67.415	68.719	69.847	71.046	71.299	72.058	65.035	65.048	63.623	62.900	65.787	66.167	65.335	67.376	63.224	61.822	61.585	60.450	61.431	62.330	61.611	65.669	62.467	
39.366	36.749	39.146	36.230	40.741	41.963	42.852	43.367	42.711	44.030	44.422	44.789	43.055	43.878	45.362	46.117	43.673	44.296	44.031	44.772	44.957	44.521	45.406	45.779	47.255	47.780	47.440	47.474	48.921	49.775	49.263	48.650	49.087	50.271	50.750	49.518	50.542	48.284	50.666	51.778	51.339
O ILE A 435 CB ILE A 435	ILE A	ILE A	11E	ASN A	ASN A	ASN A	ASN A	4	ASN A	ASN A	ASN A	ARG A	ARG A	ARG A	ARG A	ARG A	æ	ARG A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN	THR A	THR A		GLN A	GLN A									
3475	3477	3478	3479	3480	3481	3482	3483	3484	3485	3486	3487	3488	3489	3490	3491	3492	3493	3494	3495	3496	3497	3498	3499	3500	3501	3502	3503	3504	3505	3506	3507	3508	3509	3510	3511	3512	3513	3514	3515	3516

38 16.96 115.04 116.26 117.35 117.35 114.45 117.30 118.87 119.07 66.941 66.941 66.779 68.786 69.718 69.718 69.218 66.779 65.380 69.305 69 60.330 60.830 60.830 60.930 60.030 60.093 60 22.898
22.799
21.681
22.715
23.929
23.929
23.170
23.25.217
25.201
25.201
25.201
25.201
26.039
27.344
26.039
27.344
27.344
27.399
27.391
27.391
27.392
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393
27.393 35.596 35.597 35.597 36.007 44.5. 44.5. 45.5. 46.5. 47 1155. 116. 1175. 117 11.00 11 68.223
71.501
72.665
533
72.615
72.616
72.616
73.021
73.021
73.021
73.021
73.021
73.021
73.021
73.021
73.021
73.021
73.021
73.021
74.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021
75.021 63.906
64.500
65.919
66.925
66.925
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936
66.936 34.1.59 31.1.59 31.1.59 31.1.59 31.4.50 31.4.50 31.6.39 31.6.3 \* CG2 11E
CD1 11E
N SER
N SER
N SER
C SER
C SER
C SER
C SER
C C SER
N GLY
C GLY 35559 35662 35663 36663

55.984 56.808 55.525 55.427 54.854 85.400 52.620 51.967 51.967 52.620 52.620 51.967 52.6200 52.6200 52.6 48.422 44.47.546 46.813 49.689 50.149 50.149 50.13 50.148 50.148 51.420 38 0.011 38 8.011 39 8.011 39 8.016 39 8.016 30 9.018 30 9.018 30 9.013 31 9.013 31 9.013 32 9.013 33 9.013 34 9.013 36 9.013 37 9.0 15.26 115.37 115.39 116.16 116.11 116.11 117.03 117 16.44 13.04 13.89 14.71 13.39 13.08 57.374 56.424 57.023 56.369 56.171 55.149 55.284 53.743 66.241 67.497 65.325 67.497 65.028 66.028 66.128 67.128 67.128 63.247 62.731 63.857 60.410 60.410 60.228 61.521 61.521 61.528 61.528 61.528 61.528 62.431 63.837 63.837 64.41 65.62 65.62 67.638 67.63 55.532 56.514 54.662 55.165 54.262 53.387 53.933 53.174 54.297 54.070 54.070 56.211 50.211 51.326 51 51.418 50.815 51.365 50.274 50.783 50.257 49.696 51.242 50.814 49.742 49.331 52.115 28.122
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799
27.799 

																					_																				
,	oφ	80	9	8	9	æ	7	9	9	80	9	9	9	7	9	9	œ	9	7	9	9	88	9	89	. 7	9	9	80	9	9	9	9	9	9	9	7	9	9	8	9	8
	1.00 14.26		0.70 19.66			0.30 20.85		1.00 16.01	1.00 14.42	1.00 16.73	1.00 12.17					1.00 19.79								1.00 22.40				1.00 17.08	1.00 14.92	1.00 15.98			1.00 18.16			1.00 15.04	1.00 12.66	1.00 14.82	1.00 16.40	1.00 19.94	1.00 21.84
	73.451	73.358	74.900	75.257	74.932	75.127	72.188	71.766	72.699	72.913	70.320	69.872	69.389	73.236	74.181	73.501	72.298	74.731	74.181	73.501	72.855	73.329	74.569	75.047	71.674	70.881	70.092	70.019	69.938	68.846	69.087	64.289	68.092	609.99	66.848	69.565	68.696	67.223	66.885	69.002	70.360
	62.222 61.026	59.905	62.557	63.803	62.465	62.822	61.284	60.248	60.211	61.277	60.539	59.377	60.682	59.035	58.962	59.401	59.204	57.543	690.09	60.518	59.392	58.231	61.218	62.348	59.656	58.638	59.308	60.548	57.834	58.695	59.437	58.728	60.227	59.513	60.255	58.476	59.036	58.783	57.641	58.332	58.543
4	28.563	29.103	28.931	28.330	28.870	30.220	30.430	31.322	32.549	33.117	31.852	32.718	30.665	32.802	33.956	35.248	35.398	34.031	36.152	37.405	38.170	38.177	38.262	37.554	38.774	39.449	40.575	40.568	38.538	37.888	36.748	38.493	36.160	37.899	36.770	41.459	42.525	42.186	41.811	43.859	44.253
	CA SER A 476 C SER A 476	SER	CB ASER A 476		CB BSER A 476	OG BSER A 476	VAL A	CA VAL A 477	Ą	O VAL A 477	CB VAL A 477	CG1 VAL A 477	CG2 VAL A 477	4	ALA A	C ALA A 478	ALA A	ALA A	SER	SER A	SER A	SER A		SER A	PHE A	PHE A	æ	PHE A	PHE A	PHE A	PHE A	PHE A	ø	2 PHE A	K	THR	THR A	A	THR A	CB THR A 481	٠.
	3764	3766	3767	3768	3767	3768	3769	3770	3771	3772	3773	3774	3775	3776	3777	3778	3779	3780	3781	3782	3783	3784	3785	3786	3787	3788	3789	3790	3791	3792	3793	3794	3795	3796	3797	3798	3799	3800	3801	3802	3803
•	<b>ω</b> α	7	9	9	α	9	9	9	9	7	9	9	80	9	89	7	ø	φ	80	9	9	ø	1	9	9	œ	9	80	7	9	9	80	9	9	œ	7	7	9	9	89	7
1	1.00 14.55	14.		13.	19.	1.00 16.37	1.00 17.51	1.00 16.76		1.00 15.71	1.00 14.51	1.00 14.83	1.00 17.55	1.00 22.00	1.00 27.40		1.00 14.15	1.00 15.07	1.00 19.05	1.00 15.53	1.00 14.81		1.00 15.80	1.00 16.44	1.00 16.40	1.00 19.98	1.00 21.00	1.00 31.70	1.00 18.40	1.00 19.45		1.00 15.86	1.00 22.35	1.00 31.24	1.00 33.57	1.00 32.67	1.00 17.78	1.00 20.56	1.00 18.34	1.00 20.59	1.00 18.39
	67.640 68.176	67.954	68.921	69.860	69.553	68.368	67.429	67.581	67.032	70.980	71.903	72.165	72.402	73.220	74.176	72.054	72.186	73.425	73.563	70.900	70.970	69.691	74.184	75.312	75.420	75.548	76.594	77.686	75.308	75.444	74.646	75.103	76.968	77.722	17.191	78.890	73.341	72.438	72.430	71.690	73.305
	53.309	53.624	54.650	54.102	53.108	56.004	55.753	56.652	57.104	54.804	54.477	55.755	56.814	54.018	53.933	55.712	56.834	56.696	55.638	56.903	58.057	57.153	57.767	57.736	59.073	60.084	57.446	57.573	59.107	60.348	61.523	62.658	60.707	59.572	58.887	59.303	61.223	62.311	62.573	63.464	61.894
	35.261	33.976	33.623	32.518	31.867	33.155	31.948	34.319	31.315	32.408	31.268	30.497	31.107	32.024	30.967	29.172	28.298	27.380	26.764	27.433	26.446	28.376	27.351	26.386	25.711	26.435	27.214	26.284	24.396	23.632	24.194	24.363	23.685	22.991	22.086	23.579	24.452	24.866	26.369	26.807	27.146
	GLY A 469	ILE A	ILE A	ILE A	ILE A	ILE A	ILE A	ILE A	ILE A	SER A	SER A	SER A	SER A	SER A	SER A	VAL A	VAL A	VAL A	VAL A	VAL A	VAL A	VAL A	SER A	SER A	SER A	SER A	SER A	SER A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	ASN A	GLY A	GLY A	GLY A	GLY A	SER A
	00																																							0	
	3722	3724	3725	3726	3727	3728	3729	3730	3731	3732	3733	3734	3735	3736	3737	3738	3735	3740	3741	3742	3743	3744	3745	3746	3747	3748	3749	375(	3751	375	375	3754	3755	3756	3757	3758	3759	3760	3761	376	376

**		771	7.	,,,	•																																				
ω	7	9	9	œ	9	9	9	7	9	9	œ	9	9	9	9	7	9	9	9	9		9	ø	9	<b>ω</b>	ø	9	ø	œ	۲	7	9	9	œ	9	ø	9	9	9	9	9
1.00 19.75		11	12.	1.00 13.59	1.00 12.50	1.00 11.14	1.00 13.14	1.00 11.35	Η	1.00 13.16	1.00 16.40	1.00 12.61				1.00 13.76		1.00 16.82	1.00 13.91	1.00 14.61		1.00 11.75						1.00 19.79				1.00 14.21	1.00 17.30				1.00 20.95			1.00 19.51	1.00 22.52
61.907	59.626	59.340	60.269	60.410	57.882	57.650	56.882	60.914	61.883	61.508	61.324	63.283	63.638	63.904	63.716	64.166	64.049	63.546	64.216	63.701	64.044	61.394	60.903	61.637	62.207	59.410	59.117	58.993	58.485	59.455	61.523	62.092	61.102	60.614	63.393	63.984	64.638	63.800	65.176	64.324	65.003
55.439	55.975	56.313	55.457	54.276	55.903	56.154	56.812	56.135	55.413	55.715	56.878	55.953	55.807	54.628	56.884	54.969	56.328	58.271	57.117	59.052	58.461	54.618	54.869	54.121	53.092	54.452	52.972	52.219	52.726	50.952	54.698		54.203	55.341	54.774	54.219	53.010	54.959	52.511	54.444	53.254
36.431	33.950	32.567	31.685	31.910	32.231	30.712	32.978	177.08	29.900	28.446	28.103	30.222	31.708	32.381	32.632	33.717	33.883	32.509	35.020	33.641	34.894	27.680	26.332	25.221	25.472	26.259	26.541	25.208	24.222	25.214	24.024	22.819	21.665	21.507	22.452	21.152	21.146	19.963	19.956	18.770	18.817
SER A 488	Ø	K	VAL A 489	TRP A 490	TRP A 490	TRP A 490		TRP A 490	TRP A 490	ø	TRP	K	TRP	TRP A	TRP A	TRP A	TRP A	æ	ø	ø		æ	K	GLN A 491	GLN	GLN A	ø	K	<b>TYR A 492</b>			•	TYR A 492	Æ	TYR A	TYR A	TYR A 492				
8	z	క	ပ	0	ප	8	CG2	z	ర	ပ	0	CB	ខ	9	6	NEI	CE2	CE3	CZ2	CZ3	CH2	z	ð	ပ	0	8	ဗ္ဗ	8	OE1	NE2	z	ð	บ	0	8	ဗ	6	CDS	CE1	CE2	CZ
3846	3847	3848	3849	3850	3851	3852	3853	3854	3855	3856	3857	3858	3859	3860	3861	3862	3863	3864	3865	3866	3867	3868	3869	3870	3871	3872	3873	3874	3875	3876	3877	3878	3879	3880	3881	3882	3883	3884	3885	3886	3887
9	7	9	9	œ	9	9	9	9	7	9	9	8	9	7	9	9	80	9	9	9	7	9	9	80	7	9	9	80	9	7	9	9	ω	9	9	9	7	9	9	æ	9
		1.00 13.06	1.00 14.32	1.00 15.20	1.00 13.14	1.00 14.96	1.00 16.24	1.00 14.36	1.00 12.65	1.00 13.54	1.00 13.15	1.00 13.10	1.00 12.82	1.00 14.66	1.00 11.27	1.00 12.98	1.00 13.40	1.00 15.48	1.00 15.92	1.00 15.61	1.00 14.03	1.00 13.19	1.00 16.21	1.00 13.21		1.00 13.98	1.00 15.41	1.00 13.33	1.00 14.43	1.00 12.17	1.00 9.79	1.00 9.96	1.00 12.30	1.00 10.95	1.00 12.89	1.00 15.13	1.00 11.84	1.00 12.18	1.00 11.64	1.00 12.81	1.00 15.82
68.091	66.454	64.991	64.394	64.487	64.486	65.33	62.720	62.727	63.680	63.175	62.000	61.387	62.782	61.687	60.632	59.301	58.884	60.545	61.936	62.333	58.671	57.309	57.376	56.271	58.553	58.545	58.068	58.196	60.002	57.647	57.403	58.257	58.179	55.919	55.762	54.964	59.078	59.865	59.698	59.650	61.336
58.894	59.860	59.690	59.527	60.432	60.917	60.936	59.717	62.242	58.414	58.090	58.961	59.723	56.581	58.980	59.827	59.496	58.331	59.546	58.945	58.160	60.548	60.352	59.600	59.235	59.349	58.395	58.969	60.148	57.937	58.008	58.220	57.174	55.970	58.022	58.150	58.990	57.635	56.623	56.972	58.154	56.705
44.953	42.216	42.004	43.383	44.199	41.231	40.812	39.938	40.073	43.621	44.961	45.370	44.562	44.892	46.637	47.183	46.509	46.374	48.704	48.959	47.698	45.977	45.399	44.067	43.561	43.508	42.375	41.058	40.730	42.233	40.213	38.776	38.057	38.406	38.428	36.871	39.127	37.112	36.335	34.867	34.519	36.850
32 THR A 481	LEU A 482	A LEU A 482	LEU A	LEU	ALLA A	ALA A	ALA A	ALA	ALA A	PRO	PRO A	GLY A	GLY A	GLY A 485	GLY A 485	ALA A	ALA A	ALA A	ALA A 486	ALA A	VAL A	VAL A	VAL A	VAL	VAL A	VAL A	Ø	SER A	A SER A 488	SER A 488	SER	SER A									
r CG2	z			0									83									5			Z				8		S.			G						0	
3804	3805	3806	3807	3808	3809	3810	3811	3812	3813	3814	3815	3816	3817	3818	3815	3820	3821	3822	3823	3824	3825	3826	3827	3828	3829	3830	3831	3832	3833	3834	3835	3836	3837	3838	3839	3840	3841	3842	3843	3844	3845

DC7	r m	1/00	MAG	000
PC	UZ	NY:	//U\	woo

WO 99/43794

15.95 17.99 17.99 12.56 12.56 12.57 14.31 15.70 112.31 112.99 112.99 112.99 112.99 113.99 113.99 114.01 114.01 115.99 117.99 117.99 117.99 117.99 117.99 117.99 117.99 59.419
53.054
51.667
51.667
51.667
51.667
51.667
51.667
51.667
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51.675
51 44.940 59.294
59.796
59.797
60.0939
60.0939
60.0939
60.0939
60.0939
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986
60.0986 4.930 3.927 4.205 8.288 8.288 9.423 9.423 9.621 10.001 10.001 10.008 9.662 10.109 11.239 11.294 11.294 11.396 11.394 11.3 33928 33939 33939 33939 33939 33939 33946 33947 33947 33947 33947 33959 33959 33959 33960 33961 33961 33961 33961 33961 33961 33661 366 24.89
115.17
114.91
22.90
22.90
22.90
22.90
23.90
25.90
26.59
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90
27.90 65.511 60.925 60.925 60.925 60.925 60.393 60.906 60.060 60.060 60.157 60 53.781 53.151 53.151 53.353 55.462 55.462 55.291 52.191 52.191 52.191 52.191 53.429 54.967 55.339 55.339 56.039 56.030 56 17.596 20.879 19.666 18.548 19.887 20.186 17.344 14.999 14.999 15.368 15.368 115.373 115.373 117.072 1 7,226 6,242 5,049 7,183 7,578 6,685 

12.55 14.21 15.21 10.23 113.76 115.27 115.27 111.63 111.63 111.63 111.63 111.63 111.63 112.89 114.88 117.16 65.892 64.629 64.765 64.776 65.776 65.776 65.730 66.023 66.023 66.023 67.776 67 12.95 112.95 112.33 12.33 12.33 12.33 13.30 10.92 110.92 110.92 110.92 110.92 110.93 1 38.661 38.410 39.194 36.887 36.887 37.471 39.909 64.766 66.020 67.182 66.020 67.182 66.313 69.493 70.447 70.957 70.087 70 5506 5006 

92 36.63 34.71 34.71 34.71 38.71 38.71 38.73 38.73 39.73 39.73 39.73 39.73 39.74 49 .304
48 .695
49 .504
49 .595
49 .594
49 .695
49 .695
49 .696
49 .696
49 .696
49 .696
49 .696
49 .696
49 .696
49 .696
49 .696 48.808 240 47.425 51.308 51.308 52.315 51.862 51.432 51.027 50.456 51.159 52.265 51.920 647 320 657 46.155 £9. 48.485 50.076 51.530 52.864 53.867 53.965 53.965 53.965 53.965 53.965 53.965 53.965 53.965 53.965 53.965 53.965 53.965 53.965 53.965 53.965 53.965 53.965 53.965 53.965 54.518 56.975 56.975 66.984 48.984 47.505 47.020 47.251 49.673 49.948 51.367 52.313 49.835 
 14R

 14R
 14.12 13.11 14.37 17.62 17.62 17.62 18.39 14.21 13.39 14.21 13.50 15.70 17.22 16.54 17.43 17.22 17.43 17.22 17.43 20.45 117.55 115.40 117.80 20.21 116.35 114.66 23.38 26.21 27.39 26.00 24.30 26.74 1.00 1.00 1.00 1.00 1.00 1.00 1.00 54.385 54.747 54.747 55.698 56.459 57.728 55.627 54.874 55.525 53.560 52.638 53.60 52.638 51.342 51. .134 .696 .545 49.422 50.214 53.488 53.586 52.260 51.167 52.316 757 549 976 435 791 351 962 988 931 58.802 59.161 60.347 58.207 58.207 58.8136 58.8136 58.8136 58.814 58.8136 58.8 56.605 56.452 58.778 49.785 49.010 49.241 57.475 58.607 52.902 51.721 51.037 11.743 11.868 11.922 13.106 13.106 13.289 11.769 11.769 11.059 10.867 11.059 10.867 11.059 11 6.513 6.601 7.262 7.480 8.142 8.257 \*\*\* AAAAAAAAAA 

PCT/DK99/00088 WO 99/43794 15.38 119.21 118.30 118.90 118.90 118.90 118.31 119.31 110.93 110 40.489 40.679 38.998 38.998 38.998 41.415 41.415 40.613 40.613 40.613 30.7246 30.7246 31.052 34.672 42.999 43.306 44.672 44.052 45.052 46.053 46.008 46.101 47.502 48.584 48.006 49.715 49.715 47.440 50.201 559.693 559 -3.151 -1.865 -3.564 -4.6823 -3.564 -1.9237 -0.704 -0.1926 -0.1936 
 THR

 THR

 THR

 THR

 THR

 VAL

 LVS

 PRP

 TRP

 TRP

 TRP

 TRP

 TRP

 TRP

 TRP
 4180 41180 41187 41188 41188 41188 41188 41188 41189 41199 15.93 17.69 14.04 15.03 16.03 16.16 13.01 13.01 13.21 14.36 16.70 16.70 16.70 16.70 16.70 16.70 16.70 17.41 17.41 17.44 17.64 1.000 48.090
445.569
446.569
447.340
447.340
447.369
447.369
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370
447.370 66.621 66.722 67.691 68.822 70.203 70.203 70.203 61.777 66.431 66.731 67.702 72.482 70.804 71.702 72.482 70.804 70.804 70.804 70.804 70.804 70.804 70.804 70.804 70.804 70.804 70.804 70.804 70.804 70.804 66.334 66 -4.368 -5.463 -3.923 -4.592 -4.197 -3.019 -4.368 -4.368 -4.368 -4.368 1.611 1.438 2.662 2.288 3.545 3.385 0.118 0.118 -0.569 -1.992 -2.928 -3.524 -4.827 -4.271 -4.796 -6.065

4138 CB THR A 530
4140 CG2 THR A 530
4141 N PHE A 531
4142 CA PHE A 531
4143 C PHE A 531
4144 C PHE A 531
4144 C PHE A 531
4145 CB PHE A 531
4146 CG PHE A 531
4147 CD1 PHE A 531
4149 CG1 PHE A 531
4149 CG1 PHE A 531
4151 CZ PHE A 531
4151 CZ PHE A 531
4151 CZ PHE A 531
4152 C GLY A 532
4154 C GLY A 533
4155 C GLY A 533
4156 C GLY A 534
4161 CA VAL A 534
4161 CA VAL A 534
4161 CA VAL A 534
4161 CB THR A 535
4170 C THR A 535
4171 CB THR A 536
4176 C ALA A 536
4177 CB THR A 536
4178 CB ALA A 536
4178 CB ALA A 536

N THR A 542 5.297 52.107 46.615 1.00 18	52.107 4	4 4	6.615 1.00 18	1.00 18	.77	7	4260	CG2	ILE A 546	3.624	57.682	45.030	1.00 14.12	6
A 542 6.111 51.224 47.437 1.00 20.27	51.224 47.437 1.00 20.27	47.437 1.00 20.27	1.00 20.27	20.27	9	_	4261	ទី	Ø	4.473	57.773	48.119	1.00 13.44	9
A 542 6.477 51.972 48.690 1.00 21.74	51.972 48.690 1.00 21.74	48.690 1.00 21.74	1.00 21.74	21.74	•		4262	z	Z.		56.777	42.400		7
A 542 6.369 53.228 48.814 1.00 17.27	53.228 48.814 1.00 17.27	48.814 1.00 17.27	1.00 17.27	17.27	~	80	4263	ర	Ø	•	57.543	41.134		9
A 542 7.356 50.641 46.743 1.	50.641 46.743 1.00	46.743 1.00	1.00			9	4264	U ·	GLU A 547	3.943	58.116	40.945	1.00 15.30	9
A 542 8.305 51.745 46.576 1.00 21	51.745 46.576 1.00 21	46.576 1.00 21	1.00 21	21		<b>&amp;</b>	4265	0	ø	•	57.405	41.270		8
A 542 7.091 49.930 45.442 1.00 26.69	49.930 45.442 1.00 26.69	45.442 1.00 26.69	1.00 26.69	26.69	_	vo.	4266	පි	4	•	56.644	39,913		9
A 543 7.123 51.252 49.648 1.00 18.34	51.252 49.648 1.00 18.34	49.648 1.00 18.34	1.00 18.34	18.34	•	7	4267	පි	K	6.987	56.074	39.890		9
A 543 7.474 51.808 50.923 1.00 18.96	51.808 50.923 1.00 18.96	50.923 1.00 18.96	1.00 18.96	18.96	_	w	4268	8		7.105	55.058	38.781		9
A 543 8.463 52.975 50.734 1.00 17	52.975 50.734 1.00 17	50.734 1.00 17	1.00 17	13		9	4269	Œ		6.335	54.046	38.730		œ
A 543 8.525 53.808 51.615 1.00	53.808 51.615 1.00	51.615 1.00	1.00			80	4270	OE2	ø	7.924	55.225	37.834	1.00 16.92	89
A 543 8.201 50.711 51.743 1	50.711 51.743 1.00	51.743 1.00	1.00			9	4271	z	4	3.865	59.377	40.603		7
A 543 7.254 49.673 51.954 1.00 38.42	49.673 51.954 1.00 38.42	51.954 1.00 38.42	1.00 38.42	38.42		60	4272	ਰ		2.565	60.033	40.428	1.00 13.16	9
A 544 9.313 52.865 49.721 1.00 17.78	52.865 49.721 1.00 17.78	49.721 1.00 17.78	1.00 17.78	17.78	•	7	4273	ပ	Ø	2.565	60.764	39.095		9
A 544 10.349 53.917 49.575 1.00 14	53.917 49.575 1.00 14	49.575 1.00 14	1.00 14	ř	Ť	10	4274	0	Ø	3.587	61.028	38.477		89
A 544 10.208 54.723	54.723 48.287 1.00 19	48.287 1.00 1	1.00 1	ä	~	w	4275	ප	K	2.266	61.100	41.499		9
A 544 11.018 55.668 48.072 1.00 16	55.668 48.072 1.00 16	48.072 1.00 16	1.00 16	ř		80	4276	G	4	2.134	60.409	42.872		9
A 544 11.734 53.252 49.583 0.50	53.252 49.583 0.50	49.583 0.50	0.50		_	w	4277	CG2	VAL A 548	3.376	62.179	41.584		9
A 544 12.145 52.868 51.005 0.50	52.868 51.005 0.50	51.005 0.50	0.50			9	4278	z	ø	1.338	611.119	38.644		7
A 544 11.394 53.024 51.976 0.50	53.024 51.976 0.50	51.976 0.50	0.50			80	4279	రే	<b>TYR A 549</b>	1.226	61.997	37,481		9
A 544 13.359 52.364	52.364 51.118 0.50	51.118 0.50	0.50			7	4280	ပ	TYR A 549	0.902	63.397	37.975		9
A 544 11.746 53.263 49.523 0.50	53.263 49.523 0.50	49.523 0.50	0.50			9	4281	0	4	0.223	63.571	39.016	1.00 14.84	
A 544 11.998 52.552 50.860 0.50	52.552 50.860 0.50	50.860 0.50	0.50			9	4282	6	ø	0.000	61.611	36,605		4
A 544 12.195 53.205 51.884 0.50	53.205 51.884 0.50	51.884 0.50	0.50			<b>6</b>	4283	පි		0.208	60.240	36.037		9
A 544 11.914 51.250 50.767 0.50 19	51.250 50.767 0.50 19	50.767 0.50 19	0.50 19	13		7	4284	6	ø	1.049	60.058	34.934		9
A 545 9.224 54.409 47.426 1.00	54.409 47.426 1.00 14	47.426 1.00 14	1.00 14	14		7	4285	CDS	æ	-0.398	59.160	36.628	1.00 14.96	9
A 545 9.193 55.190 46.167 1.00 16	55.190 46.167 1.00 16	46.167 1.00 16	1.00 16	16		9	4286	Œ	ď	1.252	58.788	34.395		9
A 545 7.727 55.246 45.682 1.00	55.246 45.682 1.00	45.682 1.00	1.00			9	4287	CE2		-0.214	57.885	36.081	1.00 20.11	9
A 545 7.083 54.204 45.539 1.00	54.204 45.539 1.00	45.539 1.00	1.00	1.00 17.25	_	60	4288	22	TYR A 549	0.577	57.730	34.984	1.00 20.28	9
A 545 10.085 54.589 45.084 1.00	54.589 45.084 1.00	45.084 1.00	1.00	1.00 17.96	_	G	4289	HO	Ø	0.789	56.436	34.508		8
A 545 9.964 55.404 43.794 1.00 17.03	55.404 43.794 1.00 17.03	43.794 1.00 17.03	1.00 17.03	17.03	•	vo	4290	z	K	1.626	64.446	37.496		7
A 545 10.778 54.728 42.653 1.00	54.728 42.653 1.00	42.653 1.00	1.00	1.00 15.55		S	4291	ర	VAL A 550	1.317	65.794	37.957	1.00 13.30	9
A 545 12.186 54.934 43.045 1.00	54.934 43.045 1.00	43.045 1.00	1.00			7	4292	ບ	VAL A 550	-0.145	66.139	37.602	1.00 12.86	9
A 545 13.164 54.094 42.735 1.00	54.094 42.735 1.00	42.735 1.00	1.00			9	4293	0	VAL A 550	-0.589	65.893	36.503	1.00 15.42	œ
A 545	53.020 41.999 1.00 26	41.999 1.00 26	1.00 26	26		7	4294	8	VAL A 550	2.195	66.791	37,131	1.00 12.38	9
A 545 14.392 54.343 43.179 1.00 26	54.343 43.179 1.00 26	43.179 1.00 26	1.00 26	56		7	4295	CG1	VAL A 550	1.968	68.236	37.581	1.00 13.91	9
A 546 7.282 56.458 45.332 1.00 18	56.458 45.332 1.00 18	45.332 1.00 18	1.00 18	18	•	7	4296	CG2	4	3.657	66.404	37.542		9
A 546 5.908 56.613 44.793 1.00 14.23	56.613 44.793 1.00 14.23	44.793 1.00 14.23	1.00 14.23	14.23	•	S	4297	z	K	-0.828	66.685	38.603		7
A 546 6.077 57.267 43.406 1.00 17	57.267 43.406 1.00 17	43.406 1.00 17	1.00 17	17		9	4298	ర	Ø,	-2.272	66.942	38.387		9
A 546 6.771 58.296 43.389 1.00 15	58.296 43.389 1.00 15	43.389 1.00 15	1.00 15	15	_	m	4299	υ	PRO A 551	-2.447	68.081	37.424	1.00 16.09	9
A 546 5.039 57.498 45.668 1.00	57.498 45.668 1.00	45.668 1.00	1.00	1.00 13.38 (	_	10	4300	0	4	-1.599	68.955	37.236	1.00 16.43	œ
3 546 4 895 56.781 47 045 1 00	56 781 47 045 1 00	47 045 1 00	145 1 00	1 00 15.14			4301	£	PRO A 551	-2 869	711 737	19 755		
	101.00	-	******	11.01		,	1	3	;	200		1		,

95 13.53 13.26 17.40 17.40 13.77 13.77 13.77 13.88 23.88 28.55 28.55 11.28 11.28 11.28 11.58 11.58 11.59 47.465 47.392 48.826 49.616 46.555 46.488 45.122 49.097 50.269 50.269 51.206 52.656 53.312 53.312 54.033 55.0000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.0000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.0000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.0000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.0000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.0000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.0000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.0000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.0000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.0000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.0000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.000 55.00 45.741 46.379 44.044 45.010 47.33320 47.739 47.754 47.579 49.195 49.201 50.251 51.036 .672 72.927
72.402
73.3378
73.3378
742.402
76.093
772.093
772.093
772.093
772.093
772.093
773.219
773.219
773.219
773.219
773.219
774.093
775.093
776.015
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
777.093
7 4.469 2.616 1.566 1.566 4.351 3.750 4.129 4.129 4.129 4.129 4.161 5.186 5.186 5.186 5.186 5.186 5.186 7.186 2.947 2.525 2.525 1.751 1.751 2.150 3.312 4.473 4.473 0.603 0.603 1.542 1.542 1.367 20.55 117.00 117.00 117.00 118.18 119.65 119 40.707
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973
39.973 66.991
66.991
66.991
70.146
68.156
69.246
68.156
69.246
68.156
69.246
68.156
69.246
68.156
69.246
70.256
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266
70.266 4.017 4.017 5.630 5.630 6.635 6. 0.947 0.947 1.568 1.051 1.375 2.874 3.602 4.312 

																	٠.	-																	
9 9	8	φ,	9 9	9	۲.	w w	9 00	9	ø	9	9 4	ט ע	9	80	7	9	96	ο φ	æ	<b>છ</b>	ω (	٠ ٧	w	89	9	ø	9	Q	9	9	ø	ထ	~	<b>9</b> 4	9
12.49	6.79	5.18	2.26	22.04	.3.65	12.87	14.71	12.62	13.61	13.57	15.40	13.80	14.33	13.40	13.37	12.21	12.75	17.33	22.77	15.40	16.17	95.11	13.28	12.17	11.94	11.37	12.69	11.85	13.23	13.93	11.32	12.54	12.18	12.18	6.9
1.00.1			1.00.1			8.9						00.1					00.1					2 6										1.00	1.00	00.1	2
131 1		• • •		•	946	49.067	7.262	101	121	835	845	7.57	200	997	17.387													162	154		38.366	078	627	820	0
52			53.250				47			46		46.257			4	•		46.162			46.149				42.405					38.			40.	39.1	0
68.878	69.355	70.286	70.396	69.448	68.878	69.100	69.399	67.704	67.679	68.175	67.161	68.168	67.629	67.607	70.874	71.734	71.055	72.999	73.767	73.025	72.728	71 077	72.249	72.860	69.857	69.379	68.521	69.781	68.052	69.313	68.466	68.006	72.510	73.682	00000
6.705	9.024	6.147	4.911	3.834	7.747	8.930	7.347	9.480	10.887	11.079	11.946	12.361	13.400	14.710	9.060	8.571	8.665	9.436	9.459	9.408	10.793	7. 913	7.374	6.344	7.041	6.917	7.921	5.867	7.863	5.792	96.19	6.679	7.992	7.578	0
4 571 A 571			A 571 A 571		A 572	A 572		A 572	A 572			A 572					A 573	A 573	A 573		A 573								A 574		A 574	A 574	A 575	A 575	n n
LEU	LEU	LEU	LEU LEU	TEN	TYR	TYR	TYR	TYR	TYR	TYR	TYR	TYR	TYR	TYR									TYR			TYR	TYR	TYR		TYR		TYR	ASN	ASN	Ž
ర్ ర	0	8	පු පු	69	z	රි ර	ہ د	8	ខ្ល	9	9	E 6	C	Ю	z	ð	υ c	වී	8	ප	8 ;	z	5 0	0	CB						CZ	Ю	z	ජි ර	J
4428	4430	4431	4432	4434	4435	4436	4437	4439	4440	4441	4442	4443	4445	4446	4447	4448	4449	4451	4452	4452	4453	4 4 4	4455	4456	4457	4458	4459	4460	4461	4462	4463	4464	4465	4466	0 #
9 1	• •	9	<b>8</b> 49	7	9	φ	۰ ۵	9	9	œ	ر .	y y	ω .	ų	9	9	r 4	υ	æ	9	ωı	~ u	o vo	89	9	8	7	9	9	80	9	9	80		
0.36	24.26	.28	21.58 25.03	30.08	31.09	35.06	32.22	31.70	31.07	32.69	26.17	21.45	18.75	26.89	30.00	33.86	77.61	19.28	21.35	29.69	41.18	27.61	17.17	17.53	17.58	15.50	14.77	3.84	3.33	6.50	17.22	6.95	5.12	16.51	4.02
9.6		C		0			າຕ	m	c	E)	N.	α -		^1	_			4	2	7	4.	٦٠	4		-		_			_		М	٦	٦,	4
1.00			1.00 21				1 00					1.00					1.00		1.00	1.00	1.00	00.1						-	1.00 1	1.00		1.00	1.00	1.00	7.00
54.705 1.00 54.548 1.00	1.00	1.00	1.00	1.00	1.00	1.00		1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	Ä	-	⊣ .	55.503 1.00	1.00	502 1.00	1.00	025 1.00	1.00	-	Н	1.00	1.00	54.758 1.00	54.745 1.00	54.023 1.00	53.486 1.UU
	54.649 1.00	56.060 1.00	57.008 1.00 54.437 1.00	56.140 1.00	57.443 1.00	58.121 1.00	57.403 1.00	58.011 1.00	58.438 1.00	58.829 1.00	58.271 1.00	58.776 1.00	56.658 1.00	59.500 1.00	60.096 1.00	60.567 1.00	57.699 1.00	56.688 1.00	57.763 1.	56.156 1	57.266 1	55.503 1	55.568 1.00	55.502 1.00	54.137 1.00	53.025 1.00	55.538 1.00	55.253 1.00 1	53.752 1	52.996 1.00	55.435 1.00	54.758 1.	54	2, 5	ņ
61.112	57.258 54.649 1.00	56.845 56.060 1.00	57.536 57.008 1.00 56.711 54.437 1.00	55.968 56.140 1.00	55.701 57.443 1.00	56.887 58.121 1.00	57.873 57.403 1.00	59.083 58.011 1.00	60.107 58.438 1.00	61.213 58.829 1.00	59.887 58.271 1.00	60.798 58.776 1.00	60.707 56.658 1.00	60.023 59.500 1.00	60.946 60.096 1.00	59.088 60.567 1.00	62.774 57.699 1.00	63.585 56.688 1.00	63.752 57.763 1.	64.705 56.156 1	65.549 57.266 1	63.581 55.503 1	65.245 55.568 1.00	66.266 55.502 1.00	63.220 54.137 1.00	64.083 53.025 1.00	65.473 55.538 1.00	66.776 55.253 1.00 1	67.107 53.752 1	66.217 52.996 1.00	66.723 55.435 1.00	65.581 54.758 1.	64.399 54	65.853 54	66.361
	57.258 54.649 1.00	56.845 56.060 1.00	57.536 57.008 1.00 56.711 54.437 1.00	55.968 56.140 1.00	55.701 57.443 1.00	56.887 58.121 1.00	57.873 57.403 1.00	59.083 58.011 1.00	60.107 58.438 1.00	61.213 58.829 1.00	59.887 58.271 1.00	60.798 58.776 1.00	60.707 56.658 1.00	60.023 59.500 1.00	60.946 60.096 1.00	59.088 60.567 1.00	62.774 57.699 1.00	63.585 56.688 1.00	63.752 57.763 1.	64.705 56.156 1	65.549 57.266 1	63.581 55.503 1	65.245 55.568 1.00	66.266 55.502 1.00	63.220 54.137 1.00	64.083 53.025 1.00	65.473 55.538 1.00	66.776 55.253 1.00 1	67.107 53.752 1	66.217 52.996 1.00	66.723 55.435 1.00	65.581 54.758 1.	64.399 54	65.853 54	66.361
-2.281 61.112	0.245 57.258 54.649 1.00	-0.234 56.845 56.060 1.00	0.104 57,536 57.008 1.00 1.658 56,711 54.437 1.00	-1.218 55.968 56.140 1.00	-1.857 55.701 57.443 1.00	-2.488 56.887 58.121 1.00	-2.493 56.983 59.363 1.00 -3 025 57.873 57.403 1.00	-3.549 59.083 58.011 1.00	-2.523 60.107 58.438 1.00	-2.936 61.213 58.829 1.00	-1.202 59.887 58.271 1.00	-0.186 60.798 58.776 1.00	0.692 60.707 56.658 1.00	0.945 60.023 59.500 1.00	1.981 60.946 60.096 1.00	0,308 59.088 60.567 1.00	0.687 62.774 57.699 1.00	2.770 63.585 56.688 1.00	3.351 63.752 57.763 1.	0.574 64.705 56.156 1	0.595 65.549 57.266 1	3.399 63.581 55.503 L	5.229 65.245 55.568 1.00	4.519 66.266 55.502 1.00	5,381 63.220 54.137 1.00	5.066 64.083 53.025 1.00	6.572 65.473 55.538 1.00	7.143 66.776 55.253 1.00 1	6.848 67.107 53.752 1	6.527 66.217 52.996 1.00	8.670 66.723 55.435 1.00	9.363 65.581 54.758 1.	9.038 64.399 54	10.455 65.853 54	FG 185:88 BOT:/
A 563 -2.281 61.112 A 564 0 154 58 701	A 564 0.245 57.258 54.649 1.00	A 564 -0.234 56.845 56.060 1.00	A 564 0.104 57.536 57.008 1.00 A 564 1.658 56.711 54.437 1.00	A 565 -1.218 55.968 56.140 1.00	A 565 -1.857 55.701 57.443 1.00	A 565 -2.488 56.887 58.121 1.00	A 565 -2.493 56.983 59.363 1.00 A 566 -3 025 57.873 57.403 1.00	A 566 -3.549 59.083 58.011 1.00	A 566 -2.523 60.107 58.438 1.00	A 566 -2.936 61.213 58.829 1.00	A 567 -1.202 59.887 58.271 1.00	A 567 -0.186 60.798 58.776 1.00	A 567 0.692 60.707 56.658 1.00	A 567 0.945 60.023 59.500 1.00	A 567 1.981 60.946 60.096 1.00	A 567 0.308 59.088 60.567 1.00	A 568 0.687 62.774 57.699 1.00	A 568 2.770 63.585 56.688 1.00	A 568 3.351 63.752 57.763 1.	A 568 0.574 64.705 56.156 1	A 568 0.595 65.549 57.266 1	A 569 3.399 63.581 55.503 1	A 569 4.887 43.784 55.480 1.00 A 569 5.229 65.245 55.568 1.00	A 569 4.519 66.266 55.502 1.00	A 569 5.381 63.220 54.137 1.00	A 569 5.066 64.083 53.025 1.00	A 570 6.572 65.473 55.538 1.00	A 570 7.143 66.776 55.253 1.00 1	A 570 6.848 67.107 53.752 1	A 570 6.527 66.217 52.996 1.00	A 570 8.670 66.723 55.435 1.00	A 570 9.363 65.581 54.758 1.	A 570 9.038 64.399 54	A 570 10.455 65.853 54	A 571 / 108 68:381
THR A 563 -2.281 61.112	ALA A 564 0.245 57.258 54.649 1.00	ALA A 564 -0.234 56.845 56.060 1.00	ALA A 564 0.104 57,536 57.008 1.00 ALA A 564 1.658 56,711 54.437 1.00	GLY A 565 -1.218 55.968 56.140 1.00	GLY A 565 -1.857 55.701 57.443 1.00	GLY A 565 -2.488 56.887 58.121 1.00	GLY A 565 -2.493 56.983 59.363 1.00	GLY A 566 -3.549 59.083 58.011 1.00	GLY A 566 -2.523 60.107 58.438 1.00	GLY A 566 -2.936 61.213 58.829 1.00	VAL A 567 -1.202 59.887 58.271 1.00	VAL A 567 -0.186 60.798 58.776 1.00	VAL A 567 0.692 60.707 56.658 1.00	VAL A 567 0.945 60.023 59.500 1.00	VAL A 567 1.981 60.946 60.096 1.00	VAL A 567 0.308 59.088 60.567 1.00	SER A 568 0.687 62.774 57.699 1.00	SER A 568 2.770 63.585 56.688 1.00	SER A 568 3.351 63.752 57.763 1.	SER A 568 0.574 64.705 56.156 1	SER A 568 0.595 65.549 57.266 1	SER A 569 3.399 63.581 55.503 1	SER A 569 4.897 93.704 55.460 1.00 SER A 569 5.229 65.245 55.568 1.00	SER A 569 4.519 66.266 55.502 1.00	SER A 569 5.381 63.220 54.137 1.00	SER A 569 5.066 64.083 53.025 1.00	ASN A 570 6.572 65.473 55.538 1.00	ASN A 570 7.143 66.776 55.253 1.00 1	ASN A 570 6.848 67.107 53.752 1	ASN A 570 6.527 66.217 52.996 1.00	ASN A 570 8.670 66.723 55.435 1.00	ASN A 570 9.363 65.581 54.758 1.	ASN A 570 9.038 64.399 54	ASN A 570 10.455 65.853 54	LEU A 5/1 /.108 68.361 53
CG2 THR A 563 -2.281 61.112	CA ALA A 564 0.245 57.258 54.649 1.00	C ALA A 564 -0.234 56.845 56.060 1.00	A 564 0.104 57.536 57.008 1.00 A 564 1.658 56.711 54.437 1.00	N GLY A 565 -1.218 55.968 56.140 1.00	CA GLY A 565 -1.857 55.701 57.443 1.00	C GLY A 565 -2.488 56.887 58.121 1.00	0 GLY A 565 -2.493 56.983 59.363 1.00 N GLY A 566 -3 025 57 873 57 403 1.00	CA GLY A 566 -3.549 59.083 58.011 1.00	C GLY A 566 -2.523 60.107 58.438 1.00	O GLY A 566 -2.936 61.213 58.829 1.00	N VAL A 567 -1.202 59.887 58.271 1.00	CA VAL A 567 -0.186 60.798 58.776 1.00	O VAL A 567 0.692 60.707 56.658 1.00	CB VAL A 567 0.945 60.023 59.500 1.00	CG1 VAL A 567 1.981 60.946 60.096 1.00	CG2 VAL A 567 0.308 59.088 60.567 1.00	SER A 568 0.687 62.774 57.699 1.00	C SER A 568 2.770 63.585 56.688 1.00	O SER A 568 3.351 63.752 57.763 1.	CB SER A 568 0.574 64.705 56.156 1	OG SER A 568 0.595 65.549 57.266 1	N SER A 569 3.399 63.581 55.503 1	SER A 569 4.897 93.704 55.460 1.00 SER A 569 5.229 65.245 55.568 1.00	O SER A 569 4.519 66.266 55.502 1.00	SER A 569 5.381 63.220 54.137 1.00	OG SER A 569 5.066 64.083 53.025 1.00	N ASN A 570 6.572 65.473 55.538 1.00	CA ASN A 570 7.143 66.776 55.253 1.00 1	C ASN A 570 6.848 67.107 53.752 1	O ASN A 570 6.527 66.217 52.996 1.00	CB ASN A 570 8.670 66.723 55.435 1.00	CG ASN A 570 9.363 65.581 54.758 1.	OD1 ASN A 570 9.038 64.399 54	ASN A 570 10.455 65.853 54	N LEU A 5/1 /.108 68.381 53

y	o	9	8	7	7	9	9	æ	9	80	9	7	9	9	œ	9	<b>60</b>	7	ø	9	8	7	9	9	۷	9	9	æ	g	9	9	7	9	9	æ	G	G	w	w	9	9
7	ق و	ທ	4	· 		m	9	Ξ.		2	50		80	80	4		<b>ت</b>	9,	9			m	E	-		6	9	0		_ 			ص		80	_	6	و ب		و و	9
11.27	10.4		13.44	11.11				12.73		12.92		11.67												12.71						15.03			12.19	14.77	12.38	12.5	11.39	13.56	13.70	13.8	12.3
1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00
27.465	28.975	9.654	29.921	0.005	24.953	23.772	4.293	25.313	22.763	21.599	23.359	23.507	3.931	23.061	1.844	23.747	24.240	23.781	3.016	3.298	24.440	3.506	2.757	23.220	2.244	22.421	2.693	21.803	21.064	21.158	20.717	23.908	24.243	24.063	24.709	25.704	25.987	5.009	5.231	5.284	5.520
																														•	•								8 26	20	2 26
75.60	75.30	76.579	77.588	76.528	77.209	77.646	78.609	79.235	78.306	78.798	79.504	78.712	79.527	80.783	80.687	78.697	79.449	81.899	83.14	84.26	84.570	83.645	84.968	82.604	84.868	86.030	87.219	87.576	86.310	87.605	85.111	87.762	88.986	90.161	90.209	88.944	88.184	86.809	88.86	86.06	88.102
8.751	407	920	545	6.582	202	978	049	9.921	8.018	736	7.265	166	321	12.300	496	612	755	118	741	721	990	358	041	279	307	166	244	450	882	712	15.750	361	622	584	4.668	2.174	988.01	879	694	9.758	543
89	æ	7.	8.	9	89	89	10.	9.	89	ω.	7.	11.	12.	12.	12.	13.	14.	12.	Ξ.	12.	13.	10.	10.	9.	13.	14.	13.	12.	14.	15.	15.	13.	12.	13.	14.	12.	10.	10.	9.	9.	æ
581	581	581	581	581	582	582	582	582	582	582	582	583	583	583	583	583	583	584	584	584	584	584	584	584	585	585	585	585	585	585	585	286	586	586	586	586	286	586	586	586	586
K	Z,	4	ø	4	Ø	4	Ø	¥	4	4	Ø	æ	ø	¥	4	4	ø	K	¥	⋖	4	ø	Ø,	ø	4	Þ	Ø	æ	ď	æ	ø	ø	∢	K	4	₫,	ø	ø	ø,	æ	Ø
_	_	•	_	EZ GLN	THE		THR		-	OG1 THR						3 SER			-						VAL				•	•	·		A PHE	PHE		-	3 PHE	)1 PHE	_	31 PHE	CE2 PH
e C C			-		z				9 CB			z z																8							2			8 CD3	on.		
4510	451	4512	4513	4514	451	4516	451	4518	4519	4520	4521	4522	4523	4524	452	452	452	452	452	4530	453	453	453	4534	4535	453	4537	453	4539	4540	4541	454	4543	454	4545	454	4547	4548	454	4550	455
œ	g	9	æ	7	7	9	9	æ	9	9	ø	ø	7	ø	ø	æ	φ	9	9	9	7	9	9	89	9	æ	7	9	Q	<b>æ</b>	7	9	9	ထ	9	æ	9	7	9	φ	æ
12.20	. 02	00.	1.54	13.02	10.95	36.	3.05	3.39	12.95	13.10	14.18	15.07	.51	12.74	11.80	3.37	11,48	12.21	3.03	13.85	1.76	00.9	7.37	17.93	13.25	1.45	3.28	14.88	13.39	14.78	11.35	13.16	14.61	14.26	13.57	16.06	20.75	11.89	10.06		.94
			.00 14		.00		.00 13		.00	.00		.00 1	00 1	.00 13						.00				.00				.00 14		.00			.00 14	.00 14			.00 20		.00	00 10	00 13
1.00	<u>.</u> ;	4	~i	-	-	-	-	7	7	Н	H	-	-	_	Н	Н	-	٦	-	-	Н	7	Н	-	-	-	-	7	~	<b>+</b>	-4	-	-	٦	7	н	_	-	Η.	7	H
37.598	39.331	38,700	38.872	37.934	38.739	37.604	36.602	36.930	38.127	39.190	36.966	39.691	35.368	34.303	33.772	34.021	33.137	33.569	32.421	33.989	33.006	32.475	31.215	30.685	32.132	33.305	30.688	29.497	28.818	29.318	27.740	27.005	26.647	26.349	25.835	25.176	24.732	26.706	26.609	25.148	24.338
72.699	.351	.707	.292	76.223	.515	73.221	74.374	.497	73.061	71.956	72.677	71.488	.054	75.052	308	.474	74.463	74.238	.601	75.576	.385	.726	. 960	76.134	.230	.967	75.095	.270	74.010	74.450	73.227	.961	.292	75.282	600.	929.	.715	.346	75.658	. 905	.020
7.171	8.836	8.635	7.562	9.608	5.400	4.506	4.485	4.145	3.036	3.082	2.075	1.705	4.883	4.906	3.480	2.572	5.757	7.226	7.982	7.897	3.398	2.037	1.681	0.553	2.081	2.321	2.536	2.266	3.627	4.670	3.516	4.803	5.419	4.747	4.517	5.753	3.688	6.787	7.454	8.012	8.163
575						576																													580	580	580	581	581	581	581
ASN A	-			ASN A		TLE A		ILE A			ILE A				LEU A																THR A				THR A	THR A	THR A	JLN A	JLN P	JLN A	SLN ?
0																																									
4468 (																																									
44	44	44	44	44	44	44	44	44	44	44	44	44	4	44	44	44	4	44	44	44	44	44	44	44	44	44	44	44	44	44	44	45	45	45	45	45	45	45	45	45	45

98 20.17 21.50 22.70 24.71 23.44 23.44 23.19 23.27 23.29 23.23 23.39 23.39 23.39 23.39 23.39 23.39 23.39 24.57 34.82 34.82 34.82 34.82 37.82 29.550 29.779 31.094 32.102 29.941 29.941 29.941 29.941 32.102 32.292 32.292 32.292 32.292 32.292 34.303 34.318 35.620 36.918 37.918 37.918 37.918 37.918 37.918 37.918 37.918 37.918 37.918 37.918 37.918 37.918 37.918 37.918 37.435 35.407 35.681 37.092 34.628 37.473 37.288 37.949 36.544 36.399 105.367 105.252 104.537 104.537 1004.784 1006.727 1006.727 103.723 103.723 103.723 103.723 1001.521 1001.521 1001.521 1001.624 10 18.778 20.204 20.204 19.811 20.715 118.337 21.530 22.246 22.246 25.285 25.285 25.285 25.285 25.286 25.318 26.939 26.939 26.939 26.939 26.939 26.939 26.939 27.759 27.740 27.759 27.740 27.759 27.759 27.740 27.759 27.740 27.759 2 11. 70 14.16 15. 25 115. 25 115. 25 116. 20 117. 31 14. 60 117. 31 117. 31 117. 31 117. 31 117. 31 117. 31 117. 32 117. 32 117. 33 117. 34 117 26.558
23.239
23.058
23.066
23.066
23.076
23.076
24.125
25.11.118
24.125
25.076
26.231
26.831
27.125
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126
27.126 86.707
91.146
92.324
93.561
93.651
93.651
92.652
93.708
94.487
95.779
96.867
97.255
97.255
97.570
98.670
98.670
98.670
98.670
98.670
98.670
98.670
98.670
98.670
98.670
98.670 .514 .617 .871 .325 .326 .303 .529 .319 .797 .085 8.577 13.189 14.016 112.072 112.072 112.072 112.073 113.247 113.189 114.016 114.016 115.258 115.258 117.052 118.016 119.016 11 2.23 2.33

99 11.85 12.50 13.77 13.77 13.73 14.07 13.26 11.73 13.26 11.73 13.26 11.73 13.26 11.73 13.26 11.73 13.26 11.73 13.26 11.73 13.26 11.73 13.26 14.73 17.73 36.987 37.997 33.731 32.522 32.699 33.221 32.280 90.241 92.414 889.331 889.331 881.348 881.080 881.080 881.080 881.080 881.010 881.0 18.66 115.76 119 35.612 34.665 33.605 35.006 35.006 35.006 33.006 33.120 33.130 33.021 33.021 33.021 33.021 33.021 33.021 33.021 33.021 33.031 33 96.919
95.824
94.484
94.484
94.484
94.484
94.484
94.484
94.484
94.484
94.484
94.484
94.484
94.484
94.484
94.484
96.334
96.334
96.336
96.336
96.336
96.336
96.336
96.336
96.336
96.336
96.336
96.336
96.336
96.336
96.336
96.336
96.336
96.336 20 861 19 00 11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75
11.75 

100 13, 65 15, 24 12, 74 13, 71 14, 77 15, 88 16, 93 16, 93 16, 93 11, 93 11, 13 11 14.93 15.28 15.28 17.52 43.312 42.235 45.396 45.441 44.080 43.621 43.216 40.520 42.425 43.143 41.143 44.075 44.970 44.970 44.970 44.350 41.355 41.355 41.556 40.674 41.713 43.552 44.200 44.333 43.015 41.907 811 94.523 99.680 98.754 98.754 98.754 98.754 98.754 98.754 99 9.718 12.033 13.188 14.084 13.562 14.084 15.251 13.956 14.204 15.390 16.387 17.752 17.752 17.753 17. 11/162 11/162 11/163 14.30 113.57 110.078 110.078 110.078 110.078 110.078 110.078 110.088 1 39.478 38.242 38.534 39.608 40.107 38.832 37.664 35.617 35.815 34.288 34.506 33.747 40.049 41.025 33.078 31.627 30.695 31.238 36.525 38.526 39.564 39.647 39.647 39.072 40.207 39.555 38.890 37.632 36.359 39.281 41.315 42.250 41.406 41.037 36.402 86.188 85.668 85.643 85.194 86.230 86.230 86.174 86.174 86.174 87.200 88.975 88.975 88.975 88.962 88.962 88.962 88.962 89.973 89.974 89.973 89 89.943 88.719 91.648 92.394 91.869 92.464 93.904 841 572 3.501 5.010 5.298 6.284 3.915 612 613 613 613 613 614 614 614 614 LEEU GGLY Y GGLY 

101 15.67 14.76 14.76 14.70 11.21 12.14 16.93 115.73 115.92 119.58 119.58 119.58 119.59 119.58 119.59 11 18.41 18.61 28.21 27.73 19.54 28.919
30.200
28.084
28.421
27.919
26.394
27.111
27.886
27.056
27.056
27.111
27.886
27.07
26.702
26.702
26.703
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.111
27.886
27.886 21.748 19.950 19.168 18.455 22.883 21.354 19.161 91.433 91.075 92.370 91.374 91.805 93.197 90.663 89.332 89.332 96.038 94.136 95.400 95.400 95.400 95.400 95.602 97.779 96.635 97.779 96.635 97.779 96.635 97.779 97.779 96.635 97.779 97.779 98.310 100.496 100.672 99.537 100.604 99.537 100.604 100.369 101.933 101.367 102.953 22.255 23.030 23.749 23.520 22.919 16.549 15.771 15.630 20.804 20.139 20.139 22.139 22.139 23.529 23.529 24.773 24.610 22.937 24.773 24.773 25.633 25.633 27.64 27.23 27.64 27.23 27.64 27.23 27.64 27.23 27.64 27.23 27.64 27.23 27.64 27.23 27.64 27.23 27.64 27.23 27.64 27.23 27.64 27.23 27.64 27.23 2 4874 4875 4876 1877 1878 1879 1880 1882 1883 1884 1885 1885 10.99 111.13 13.90 113.90 113.90 113.90 113.91 112.44 112.64 112.64 112.70 113.72 113.72 113.73 114.06 115.64 116.93 117.93 117.93 117.93 117.94 117.93 117. 1.00 34.071 33.762 33.521 34.084 34.096 36.040 33.941 .131 .510 .003 36.947 37.469 38.120 35.631 019 320 858 .732 .084 .141 .596 .048 .322 .037 .960 41.151 40.044 42.196 39.098 37.812 34.660 34.310 934 332 881 86.477 85.492 84.323 83.546 84.078 91.012 89.927 88.780 91.061 91.116 91.609 92.215 85.753 87.626 88.339 87.592 89.748 87.740 87.034 87.912 89.092 87.264 87.950 87.91786.888 88.815 90.097 89.998 88.992 81.346 79.972 79.689 79.105 84.199 85.623 86.389 86.389 85.956 85.956 85.956 87.018 8.656 8.656 9.398 9.398 11.110 11.110 11.110 11.110 11.110 11.110 11.100 

TYR A 632 22.814 103.670 28.369 1.00 23.27 8 PRO A 633 21.113 101.067 23.157 1.00 19.06 7 PRO A 633 20.194 101.103 24.225 1.00 16.58 6	103.670 28.369 1.00 23.27 8 4930 CA TYR A 101.067 23.157 1.00 19.06 7 4931 C TYR A 101.103 24.225 1.00 16.58 6 4932 O TYR R	570 28.369 1.00 23.27 8 4930 CA TYR A 267 23.157 1.00 19.06 7 4931 C TYR A 103 24.225 1.00 16.58 6 4932 O TYR R	23.27     8     4930     CA     TYR A       19.06     7     4931     C     TYR A       16.58     6     4932     O     TYR R	4930 CA TYR A 4931 C TYR A 4932 O TYR A	CA TYR A C TYR A O TYR A	TYR A TYR A TYR A	A 63 A 63 A 63		17.759 16.714 15.911	87.841 86.848 87.135	26.425 25.989 25.088	22.22	ه <b>و</b> ، و،
A 633 19.087 100.033 24.123 1.00 15.75 6 4933 CB TYR	100.033 24.123 1.00 15.76 6 4933 CB TYR	1033 24.123 1.00 15.76 6 4933 CB TYR 15.3 24.123 1.00 15.74 8 4934 CG TYR	15.76 6 4933 CB TYR 16.24 R 4934 CG TYR	4934 CG TYR	CB TYR	TYR		A 637	18.183	86.513			o vo vo
PRO A 633 19.474 102.498 24.311 1.00 19.94 6 4935	102.498 24.311 1.00 19.94 6 4935 CD1	.498 24.311 1.00 19.94 6 4935 CD1	19.94 6 4935 CD1	4935 CD1	8 6		202	. ∢	20.253	86.244	27.548	12.	9
PRO A 633 19,449 102.810 22.846 1.00 24.00 6 4936 CD2 1 PPO A 633 20,772 102.271 22.369 1.00 24.76 6 4937 CB1	102.810 22.846 1.00 24.00 6 4936 CD2	.810 22.846 1.00 24.00 6 4936 CD2 .271 22.369 1.00 24.76 6 4937 CE1	24.00 6 4936 CD2 24.76 6 4937 CE1	4936 CD2	GE 17		TYR TYR	A 637	18.595	85.573	29.127	1.00 10.55	o o
ASP A 634 19.272 99.094 23.169 1.00 16.18 7 4938	99.094 23.169 1.00 16.18 7 4938	094 23.169 1.00 16.18 7 4938	16.18 7 4938	4938		CE2	TYR	4	19.321	84.424	29.387		9
ASP A 634 18,330 97.978 23.121 1.00 16.48 6 4939	97.978 23.121 1.00 16.48 6 4939	23.121 1.00 16.48 6 4939	16.48 6 4939	4939		Z	TYR	æ	20.521		28.728	1.00 11.69	9
ASP A 634 18,923	96.792 23.937 1.00 15.97 6 4940	23.937 1.00 15.97 6 4940	15.97 6 4940			H 2	TYR	A 637	21.255	83.073	29.016	1.00 11.85	ם ר
ASP A 634 20.153 97.402 21.731 1.00 15.05 6 4942	97.402 21.731 1.00 15.05 6 4942	21.731 1.00 15.05 6 4942	15.05 6 4942			: రో	<u> </u>	<b>A</b>	16.033	84.508	25.827		. 49
ASP A 634 17.626 98.434 20.695 1.00 20.07 6 4943	98.434 20.695 1.00 20.07 6 4943	20.695 1.00 20.07 6 4943	20.07 6 4943			υ	VAL	ø	15.492	83.693	27.023		9
ASP A 634 17.138 99.519 21.100 1.00 17.34 8	99.519 21.100 1.00 17.34 8 4944	21.100 1.00 17.34 8 4944	17.34 8 4944			0	VAL		16.202	83.367	27.980	1.00 13.35	Ф
ASP A 634 17.728 98.097 19.503 1.00 19.32 8 4945	98.097 19.503 1.00 19.32 8 4945	19.503 1.00 19.32 8 4945	19.32 8 4945			8		ø	16.830	83.496	24.987		9
TRP A 635 18.009 96.147 24.656 1.00 14.45 7 4946	96.147 24.656 1.00 14.45 7 4946	24.656 1.00 14.45 7 4946	14.45 7 4946			8		<b>4</b>	15.922	82.650	24.118	1.00 12.74	<b>ب</b> م
TRP A 635 18.413 95.017 25.510 1.00 16.15 6 4947	95.017 25.510 1.00 16.15 6 4947	25.510 1.00 16.15 6 4947 25.082 1.00 13.84 6 4948	16.15 6 4947			ÿ 2	2 VAL	A 638	17.799	84.300	24.074	1.00 10.59	9 1
TRP A 635 16.590 93.820 24.606 1.00 13.90 8 4949	93.820 24.606 1.00 13.90 8 4949	24.606 1.00 13.90 8 4949	13.90 8 4949			: ບັ		: 4	13.441	82.960	28.082	1.00 9.34	. 0
TRP A 635 18.024 95.299 26.962 1.00 15.52 6 4950	95.299 26.962 1.00 15.52 6 4950	26.962 1.00 15.52 6 4950	15.52 6 4950			ပ		ø	12.466	81.877	27.665		9
TRP A 635 18.818 96.421 27.587 1.00 14.07 6 4951	96.421 27.587 1.00 14.07 6 4951	27.587 1.00 14.07 6 4951	14.07 6 4951			0		¥	11.814	81.982	26.632	1.00 12.26	10
TRP A 635 18.737 97.739 27.178 1.00 14.13 6	97.739 27.178 1.00 14.13 6 4952	27.178 1.00 14.13 6 4952	14.13 6 4952			8	PHE		12.612	84.072	28.807	1.00 11.50	)2
TRP A 635 19:713 96:40/ 28:694 1:00 16:1/ 6 4953 TRP A 635 19:561 98:533 27:989 1:00 15:61 7 4954	96.40/ 28.634 1.00 16.1/ 6 4953 98.533 27.989 1.00 15.61 7 4954	28.634 1.00 16.17 6 4953 27.989 1.00 15.61 7 4954	15.61 7 4954			ว์ บี		4 4	14.294	85.209	30.446		و م
TRP A 635 20.179 97.706 28.908 1.00 17.77 6 4955	97.706 28.908 1.00 17.77 6 4955	28.908 1.00 17.77 6 4955	17.77 6 4955			Ü			13.567	86.380	28.563		9
TRP A 635 20.179 95.350 29.511 1.00 18.51 6 4956	95.350 29.511 1.00 18.51 6 4956	29.511 1.00 18.51 6 4956	18.51 6 4956			C		K	15.176	86.094	30.829	1.00 12.05	9
TRP A 635 21.071 98.018 29.935 1.00 19.26 6 4957	98.018 29.935 1.00 19.26 6 4957	29.935 1.00 19.26 6 4957	19.26 6 4957			υ		Ą	14.416	87.425	28.965		9
TRP A 635 21.100 95.671 30.515 1.00 23.17 6 4958	95.671 30.515 1.00 23.17 6 4958	30.515 1.00 23.17 6 4958	23.17 6 4958			O	<b>6</b> 1	ø	15.196	87.301	30.113		9
TRP A 635 21.514 96.992 30.735 1.00 20.92 6 4959	96.992 30.735 1.00 20.92 6 4959	30.735 1.00 20.92 6 4959	20.92 6 4959			z	N SER	A	12.323	80.913	28.609	1.00 10.61	7
PHE A 636 18.294 92.561 25.372 1.00 12.41 7 4960	92.561 25.372 1.00 12.41 7 4960	25.372 1.00 12.41 7 4960	12.41 7 4960			v	4	⋖	11.376	79.833	28.360		9
PHE A 636 17.562 91.335 24.947 1.00 11.57 6 4961	91.335 24.947 1.00 11.57 6 4961	24.947 1.00 11.57 6 4961	11.57 6 4961			O		æ	10.005	80.223	28.872		9
PHE A 636 18.071 90.194 25.855 1.00 12.41 6 4962	90.194 25.855 1.00 12.41 6 4962	25.855 1.00 12.41 6 4962	12.41 6 4962			0		Ø	9.878	80.649	30.026		80
PHE A 636 19.204 90.239 26.345 1.00 13.24 8 4963	90.239 26.345 1.00 13.24 8 4963	26.345 1.00 13.24 8 4963	13.24 8 4963			ប	3 SER	A 640	11.915	78.568	29.132	1.00 12.23	9
A 636 17.761 90.971 23.435 1.00 12.03 6 4964	90.971 23.435 1.00 12.03 6 4964	971 23.435 1.00 12.03 6 4964	12.03 6 4964			8	SER	A 640	11.028	77.448	28.846		æ
PHE A 636 19,030 90.118 23.261 1.00 12.15 6 4965	90.118 23.261 1.00 12.15 6 4965	118 23.261 1.00 12.15 6 4965	12.15 6 4965			z	VAL	A 641	9.007	80.190	27.975		7
. PHE A 636 20.287 90.667 23.292 1.00 17	90.667 23.292 1.00 17.38 6 4966	667 23.292 1.00 17.38 6 4966	17.38 6 4966			ర	VAL	A 641	7.635	80.609	28.374	1.00 11.37	9
PHE A 636 18.856 88.751 23.095 1.00 11.02 6 4967	88.751 23.095 1.00 11.02 6 4967	23.095 1.00 11.02 6 4967	11.02 6 4967			ပ	VAL	A 641	6.697	79.601	27.775	1.00 11.35	9
PHE A 636 21.417 89.860 23.207 1.00 19.56 6 4968	89.860 23.207 1.00 19.56 6 4968	23.207 1.00 19.56 6 4968	19.56 6 4968			0		ø	7.075	78.840	26.864	1.00 10.31	80
A 636 19.996 87.906 22.997 1.00 12.49 6 4969	87.906 22.997 1.00 12.49 6 4969	22.997 1.00 12.49 6 4969	12.49 6 4969			ü		æ	7.286	82.019	27.760	_	9
PHE A 636 21.250 88.466 23.034 1.00 17.21 6 4970	88.466 23.034 1.00 17.21 6 4970	23.034 1.00 17.21 6 4970	17.21 6 4970			ဗ	1 VAL	A 641	8.061	83.038	28.624	1.00 13.20	y
TYR A 637 17.229 89.170 25.966 1.00 12.09 7 4971	89.170 25.966 1.00 12.09 7 4971	25.966 1.00 12.09 7 4971	12.09 7 4971			U	CG2 VAL	A 641	7.607	82.115	26.241	1.00 10.82	9

103 2188011222227934444111988919892234444740468 30.490 31.338 30.611 29.820 30.260 30.260 30.260 30.155 30.155 30.051 29.862 30.051 29.863 30.051 20.633 31.338 31.405 31.338 31.328 32.306 33.318 33.328 33 564 453 426 426 723 704 31. 32. 33. 33. 33. 33. 90.407
90.407
91.407
92.407
93.608
93.608
93.608
93.708
93.708
93.708
93.708
93.708
93.708
93.708
93.708
93.708
93.708
93.708
93.708
93.708
94.708
95.708
96.708
97.708
97.708
97.708 2.687 3.041 0.298 -1.032 -2.040 3.446 4.767 5.219 4.996 6.812 5.031 6.057 6.057 6.057 9.090 9.172 10.217 11.465 11.265 11 50114 50115 50116 50116 50117 99799999779 10.92 11.56 11.56 11.56 11.56 11.37 11.37 11.37 11.34 11.37 28.254 27.626 25.888 28.529 29.308 25.291 23.839 25.291 23.141 23.141 23.141 24.236 24.236 25.291 25.291 26.493 27.269 28.279 279 279 279 279 79.384 78.438 79.384 79.994 77.999 77.999 77.999 77.999 77.999 77.999 77.999 77.999 77.999 77.999 77.999 77.999 77.999 77.999 88.109 88 5.470 4.582 4.295 3.295 

104 33.22 24.748 25.70 26.81 27.05 2 42.706 41.909 40.758 39.919 41.207 41.207 41.221 38.624 37.743 36.669 35.730 35.730 35.730 35.730 37.743 37 44.643 44.087 44.267 44.267 44.227 44.227 44.227 43.931 43.566 43.215 41.985 41.118 42.717 43.510 99.783
99.906
102.050
102.050
103.920
98.958
98.181
98.837
98.126
100.106
100.108
99.778
100.279
100.293
101.011.01.014
100.405
100.277
100.205
100.405
100.405
100.926
100.926
100.926
100.926
100.926
100.926
100.926
100.926
100.926
100.926
100.926
100.926 16.419 16.614 17.524 15.276 14.319 12.353 12.186 12.458 13.012 11.704 14.022 11.704 11.720 12.266 13.410 12.611 13.541 13.279 14.578 9.556 9.441 9.412 8.152 7.960 7.960 7.543 20.202 20.202 21.725 20.499 19.665 20.239 20.019 18.874 17.543 16.531 17.502 16.313 15.641 50098 510098 15.14 16.77 17.74 17.90 17.90 17.90 17.19 18.25 19.59 36.084
336.789
337.566.084
338.7.566.089
338.7.566.089
338.7.566.099
338.7.566.099
338.7.566.099
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.567
338.7.57
338.7.57
338.7.57
338.7.57
338.7.57
338.7.57
338.7.57
338.7.57
338.7.57
338.7.77
338.7.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
338.77
3 96.950 95.688 94.518 94.518 93.407 93.343 98.394 98.312 98.312 98.3110 98.3110 97.524 97.524 97.524 97.524 97.524 97.524 97.528 97.382 11. 634 12. 097 13. 633 11. 326 11. 326 11. 326 11. 363 11. 326 12. 097 12. 000 12. 000 12. 000 13. 633 14. 244 15. 705 16. 168 17. 705 19. 571 17. 709 18. 152 18. 152 19. 706 19. 70 
 PHE

 PARG

 ARG

 ARG

 ALA

105 15.66 116.10 116.11 116.12 117.00 114.13 117.00 26.930 26.856 27.847 26.519 27.847 26.519 27.447 26.519 27.472 28.734 28 94.747 93.408 94.431 95.666 96.4831 96.188 97.464 97.464 97.305 97.30 5182 5183 5184 5184 5186 5187 5188 5189 5189 5199 21. 46 118.92 118.92 118.93 118.93 118.93 118.93 118.64 119.74 110.95 119.75 11 39.608 38.713 38.713 39.6295 39.6295 39.713 39.740 98.218
97.223
96.960
97.223
96.960
97.223
96.961
100.250
100.857
101.517
100.973
99.526
98.503
99.581
103.085
100.710
100.839
99.581
98.403
99.581
98.403
99.581
99.440
100.438
99.581
99.581
99.440
99.581
99.581
99.581 7.781 7.444 7.7444 7.7444 6.798 8.551 7.240 8.551 8.065 8.065 8.065 8.065 8.065 8.159 8.065 8.159 8.063 8.159 8.159 8.159 8.161 8.159 8.169 8.16 5140 51410 514

106 23.218 20.334 20.334 20.334 20.336 20.616 119.929 118.765 22.256 22.256 22.256 22.256 23.306 24.469 22.366 22.366 22.366 22.366 22.366 22.366 22.366 22.366 22.366 22.366 22.366 23.308 24.469 22.366 22.366 22.366 23.368 24.469 26.769 26.769 26.705 27.316 28.317.706 28.317.706 28.317.706 28.317.706 28.317.706 28.317.706 28.317.706 28.317.706 28.317.706 28.317.706 28.317.706 28.317.706 28.317.706 28.317.706 28.317.706 28.317.706 86.867 91.378 92.583 92.781 93.451 93.751 94.052 94.052 94.052 96.732 96.732 96.732 96.732 96.732 96.732 97.7322 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.7322 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.7322 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.7322 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.7322 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.7322 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.7322 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.7322 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.7322 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.7322 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.7322 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.732 97.7322 97.732 97.7 6.611 9.363 10.236 9.280 11.807 11.642 9.639 9.639 10.238 8.730 9.639 10.245 11.284 11.245 11.376 11 552666 55267 55267 55267 55267 5527 5527 5527 5527 5528 5527 5527 5527 5528 5527 5528 5529 16.36 19.04 19.09 19.09 19.09 19.08 18. 724 19. 033 118. 803 118. 803 119. 803 119. 293 119. 293 119. 293 119. 293 119. 293 119. 043 119. 78.885 77.518 77.518 77.518 77.053 77.053 77.053 77.053 77.053 77.053 77.053 77.053 88.003 10.419 10.845 11.505 11.611 10.614 10.614 10.614 10.614 10.614 10.614 10.949 10.949 10.949 10.949 10.949 11.857 11 4.164 3.803 3.803 4.915 2.613 2.997 1.409 1.409 5.953 7.035 8.164 7.915 9.349 

9.18 110.73 111.48 111.48 111.48 111.03 10.33 10.33 10.33 111.04 111.05 110.38 28. 829 28. 292 29. 262 29. 262 29. 264 29. 448 20. 595 20. 59 73.167
71.601
72.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300
77.300 55.550 55 18.78 20.58 21.77 21.77 22.87 22.87 24.12 25.76 27.47 27.46 27 26.995
26.110
28.261
27.906
28.836
29.836
27.906
27.906
27.906
27.907
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
38.228
39.228
39.228
39.228
39.228
39.228
39.228
39.228
39.228
39.228
39.228
39.228
39.228
39.228 1000 283 99 604 1000 246 1100 246 1106 6954 1106 6954 1106 6954 1106 896 1106 896 1106 896 1106 896 1106 896 1106 896 1107 896 11 3.960 8.037 7.489 9.037 10.346 111.360 10.376 33.596 34.036 35.915 36.915 37.596 37.596 37.596 37.596 37.596 37.596 37.596 37.596 37.596 37.596 37.596 37.596 37.596 37.596 37.596 37.596 37.69 

WO 99/43794	PCT/DK99/00088
-------------	----------------

108 11. 54 111. 77 111. 70 111. 54 111. 54 111. 91 112. 08 112. 08 113. 09 113. 29 113. 29 113. 29 113. 29 113. 29 22.854 14.127 13.769 12.769 12.769 13 62.658 63.919 60.656 60.656 60.656 61.402 64.250 68.2331 64.250 68.673 69.399 60.993 6 26.088 37.981 40.945 40 14.01 196.48 198.54 118.54 117.24 117.24 117.24 117.24 117.24 117.24 118.54 118 70.364 70.312 71.238 68.523 68.523 67.561 66.433 68.304 67.478 70.263 71.468 72.412 72.412 70.263 71.468 72.412 72 71.291
70.667
70.667
70.667
70.621
70.621
70.681
71.638
71.638
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820
72.820 33. 269 34. 569 35. 569 36. 004 37. 561 37. **AAAAAAAAAAAAAAAAAAAAAAAAAAAAA** 

15.84 16.12 16.13 16.13 16.13 16.13 16.32 16.32 16.32 16.32 16.32 16.32 17.23 17.24 17.24 17.24 17.24 17.25 17.25 17.25 17.25 17.27 41.510
335.990
346.1519
347.991
348.1510
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519
349.1519 58.009 71.623 71.623 882.474 882.474 880.691 880.910 890.900 75.000 691.238 881.238 14, 144 44, 010 25, 010 1, 291 1, WAT

WWAT

W 18.284 53.938 33.829 30.252 43.636 43.606 43.606 43.606 43.606 6.013 31.215 20.822 22.815 11.992 6.013 31.215 31.215 42.264 42.264 6.013 31.215 31.215 42.264 31.215 32.334 33.33 52.273
647.305
688.357
68.357
68.357
68.357
68.357
68.928
68.928
68.928
68.713
69.928
68.713
69.928
69.728
69.738
77.662
77.662
77.662
77.662
77.662
77.662
77.662
77.662
77.662
77.662
77.662
77.662
77.662
77.662
77.662
77.662
77.662
77.662
77.662
77.662 37.048 39.032 39 \$5516

9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296
9.296 69.204
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.553
91.753
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.733
91.7333
91.7333 43.898 117.503 22.0119 22.0119 22.0119 22.0119 23.8222 23.8222 23.8222 23.8222 23.8222 23.8222 23.8222 23.8222 24.8222 25.8222 26.8 31.635 33.559 48.1159 33.550 33.550 33.550 33.550 33.550 33.550 33.550 33.1586 44.1116 41.1150 45.173 46.173 47.116 48.688 48.6866 48.686 48. 79.962 76.428 83.1173 84.156 85.406 85.406 87.008 8 014 811 15.832 335.482 316.532 317.532 317.532 317.532 317.532 317.632 > > > 5598 5500

<b>8 8</b>	ထေ	αο α	, ω	89	co c	o ac	æ	80	80	œ	<b>ω</b> σ	0 00	, α	œ	89	11	1	<b>2</b> 00	82	60	&	co co	80	œ	æ	<b>&amp;</b>	æ .	20 .	eo (	m (	æ	ao a	
23.	1.00 23.57 1.00 24.20	1.00 23.77	24.	24	1.00 24.19			1.00 24.31			1.00 24.18	1.00 24.65		1.00 24.73				1.00 24.37			1.00 24.82	1.00 25.15		1.00 25.20		1.00 25.43						1.00 25.40	1.00 25.51
18.325	42.902	51.124	32.939	53.507	23.573	16.052				29.872	18.858	39.112	11.440	46.357	17.334	56.879	22.319	53.380	53.486	52.493	7.698	39.623	19.817	47.663	49.110	42.326	22.790	37.221	23.179	54.469	56.789	18.448	50.293
62.142 56.925	53.127 78.220	57.105	106.639	66.715	47.324	77.638	50.730	54.776	74.449	72.335	77.051	49.306	55.968	84.809	50.984	65.666	72.661	58.393	52.674	64.827	61.118	80.899	45.139	84.191	71.857	57.278	93.061	82.782	95.706	70.499	68.339	91.019	70.892
13.480	7.117	49.295	17.824	15.542	27.050	10.356	24.405	15.639	-5.196	40.985	48.449	45.814	44.725	20.058	18.079	17.020	44.682	50.222	44.639	49.725	39.542	41.190	25.053	16.135	11.381	-4.512	3.805	34.832	3.711	20.209	11.697	7.254	17.078
WAT V	OWO WAT V 284 OWO WAT V 285	OWO WAT V 286	WAT V	WAT V	OWO WAT V 290	WAT V	> >	WAT V	OWO WAT V 305	WAT V	WAT V	WAT V	OWO WAT V 310	WAT V	WAT V	WAT V	WAT	WAT	WAT V	WAT	WAT V	WAT V	OWO WAT V 321	_									
5724 5725	5726 5727	5728	5730	5731	5732	5734	5735	5736	5737	5738	5739	5741	5742	5743	5744	5745	5746	5748	5749	5750	5751	5752	5754	5755	5756	5757	5758	5759	5760	5761	5762	5763	5765
<b>ω ω</b>	<b>co</b> co	œ o	ο α	89	α α	တ	60	89	Ф	œ	<b>a</b> c (	οα	ω (	ω	ထ	ထ	ω (	<b>20</b> 00	89	8	ω (	<b>20 00</b>	60	œ	80	ω .	ω .	<b>x</b>	œ ·	<b>co</b> (	ω	œ °	<b>.</b>
	22.04	22.42	22.90		23.10							22.63					מי	0 23.17	00 22.95	"		1.00 23.45			(1)					23.04	0 23.27	1.00 23.30	00 23.63
નં નં	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	1.00	2 6	1.00	1.00	1.00	1.00	1.00	1.00	1.00	Ä	٠,	-i -i	i,	ij	1.00	1.00	1.00	1.00	1.00	1.00	1.00	, i	i H
	20.742 1.00 29.447 1.00	55.081 1.00			72.156 1.00							44.099 I.U(						38.475 1.0		55.523 1.		75.008 1.							~ .	13.742 1.00	_	976	46.773 1.
48.609 51.930 1 94.552 38.603 1	68.970 20.742 85.642 29.447	49.542 55.081 1	43.276 33.862	86.290 16.425	60.466 72.156	65.837 24.376	72.803 60.509	91.014 44.772	98.395 35.969	68.974 53.447	66.167 37.375	72 507 16 552	56.257 16.556	82.268 13.713	49.792 60.562	61.061 10.006	70.834 12.781	93.829 38.475	101.476 27.805	54.281 55.523	64.024 50.895	48.607 29.481 63.595 75.008	43.626 21.918	85.153 37.181	71.419 66.331	79.697 18.591	43.105 36.443	65.021 59.364	69.710 39.367 1	86.458 13.742 1	86.883 37.755 ]	81.157 29.976	55.293 46.
51.930 1 38.603 1	68.970 20.742 85.642 29.447	49.542 55.081 1	43.276 33.862	86.290 16.425	60.466 72.156	65.837 24.376	72.803 60.509	91.014 44.772	98.395 35.969	68.974 53.447	66.167 37.375	72 507 16 552	56.257 16.556	82.268 13.713	49.792 60.562	61.061 10.006	70.834 12.781	93.829 38.475	101.476 27.805	54.281 55.523	64.024 50.895	48.607 29.481 63.595 75.008	43.626 21.918	85.153 37.181	71.419 66.331	79.697 18.591	43.105 36.443	65.021 59.364	69.710 39.367 1	86.458 13.742 1	86.883 37.755 ]	81.157 29.976	55.293 46.

112 34.878 34.878 34.878 36.029 37.94 38.049 49.068 44.166 74.430 99.282 99.582 99.582 75.038 77.038 77.038 77.038 77.080 43.193 4.3.193 4.5.193 4.5.193 4.6.20 4.6.30 4.6.30 4.6.30 4.6.30 4.6.30 4.7.30 4.6.30 4.7 5808 5809 5809 5810 5811 25.53 25.64 26.25 26.27 26.28 26.28 26.29 26.20 26.20 26.20 26.20 26.20 26.20 26.20 26.20 26.20 26.20 26.20 26.20 26.20 26.21 27.29 27.29 27.20 4.552 55.164 55.164 55.164 55.164 55.164 56.165 56.166 56. 60.915
71.214
96.310
96.310
70.224
64.9230
70.224
70.224
71.401
81.627
71.401
71.401
71.632
71.401
71.401
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632
71.632 49.200 31.661 -2.062 39.125 48.433 35.352 34.557 23.250 23.26 44.958 44. > WAT
WAT
WAT
WAT
WAT
WAT WAT WAT WAT WAT
WAT
WAT
WAT
WAT
WAT
WAT
WAT
WAT WAT 5.166 5.167 5.167 6.168

WO 99/43794	PCT/DK99/00088

113 31.54 31.71 30.69 31.38 30.51 30.90 31.36 31.36 31.36 31.36 31.36 31.37 31.37 31.35 11.551 41.178 37.234 36.134 49.717 42.768 53.771 14.882 70.095 70.095 71 4.445 7.578 35.219 36.237 54.586 56.245 38.501 31.829 75.573 16.251 52.856 9.218 2.001 23.792 42.809 25.885 42.193 20.484 7.006 46.702 39.931 84.645 70.200 70.200 85.464 665.098 48.993 70.509 83.732 84.031 84.031 84.031 84.031 84.031 84.031 84.031 84.031 84.031 84.031 86.031 8 68.458 105.053 93.867 68.664 58.438 85.349 56.769 42.137 22.333 40.228 1.666 14.531 26.307 51.579 16.514 23.770 50.377 15.979 8.486 17.712 45.187 23.220 23.220 43.353 0.954 8.998 47.484 25.632 27.887 10.456 25.474 25.474 10.456 11.666 17.99 21.000 37.163 41.761 27.194 39.487 30.446 -2.258 46.873 18.407 480 482 482 483 485 485 485 WAT WAT WAT WAT WAT WAT
WAT
WAT
WAT
WAT
WAT
WAT
WAT
WAT WAT
WAT
WAT
WAT
WAT
WAT
WAT WAT WAT WAT 5918 5919 5920 5921 5922 5923 5924 5925 5926 5927 28.21 28.81 29.13 29.13 29.43 29.43 29.13 29.34 29.13 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 1.00 21.116 34.445 34.445 45.763 48.593 48.593 48.593 65.360 19.020 11.952 11.953 11 21.940 23.996 18.498 38.760 57.406 58.879 58.664 16.010 .468 22 66.686 103.267 45.553 42.994 82.283 79.313 78.316 72.460 63.547 65.55 55.163 82.697 45.171 65.149 74.061 46.218 89.697 99.697 86.519 90.485 67.365 102.838 67.009 44.407 89.765 56.984 74.847 76.028 55.405 55.405 58.042 47.66189.316 40.619 58.076 40.919 7.439 27.438 32.690 8.726 9.726 9.880 10.831 10.831 119.126 117.933 44.723 28.389 45.865 117.844 117.833 44.723 47.723 4 -1.685 35.326 -5.217 22.655 9.970 22.156 27.456 7.899 30.182 36.835 24.378 441 4443 4444 446 WAT WAT WAT
WAT
WAT
WAT WAT WAT WAT WAT WAT WAT
WAT
WAT
WAT
WAT
WAT
WAT
WAT
WAT
WAT WAT WAT WAT WAT WAT 5.885.0 5.8

WO 99/43794	PCT/DK99/00088
-------------	----------------

31.57 8 5981 OWO WAT V 539 20.256
32.02 8 5980 OWO WAT V 31.57 8 5981 OWO WAT V
31.57 8 5980 OND MAI V
31.57 8 5981
31.57
1.00
1

																					1	1,5	5																		
ω .	œ	Œ	80	80	80	80	80	80	æ	œ	<b>c</b> o	œ	ω	80	80	æ	80	8	8	œ				æ	αο	œ	8	8	œ	ω	œ	œ	80	8	ထ	œ	80	œ	œ	œ	80
1.00 37.09	1.00 36.47				1.00 35.94			1.00 37.10			1.00 37.12	1.00 37.94		1.00 36.82		1.00 36.97		1.00 36.23				1.00 36.17	1.00 37.66	1.00 37.35	1.00 37.56				1.00 36.99	1.00 37.37			1.00 37.57	1.00 37.37	1.00 37.91	1.00 36.73	1.00 37.70	1.00 38.56		1.00 37.83	1.00 37.72
52.034	58.223	42.499	21.447	35.973	57.692	30.332	18.637	59.420	50.091	37.499	41.392	13.060	7.515	18.395	70.896	28.706	64.130	25.141	69.041	48.034	17.015	43.853	46.103	70.610	25.786	33.216	30.305	12.899	41.877	65.800	15.071	48.688	38.409	56.844	18.192	51.941	32.915	26.870	11.597	45.035	25.195
80.214	72.793	90.473	73.674	104.300	78.729	94.866	79.736	55.701	47.227	52.302	50.088	80.815	62.973	57.372	55.821	93.998	54.704	105.878	67.557	88.594	42.321	84.033	51.035	72.978	69.664	107.231	98.361	84.394	52.936	49.377	44.914	49.772	51.717	54.692	90.646	55.043	98.451	98.198	49.796	55.242	43.210
40.907	47.006	20.221	100		30.691	-3.070	46.455	46.537	43.765	-0.451	47.301			13.837									10.886					34.857	20.425	26.575	31.193	10.578	18.128	45.424	28.011	47.536	24.853	27.088	33.663	16.309	20.804
_	WAT V	WAT V	>	OWO WAT V 622	OWO WAT V 623	OWO WAT V 624	OWO WAT V 625	OWO WAT V 626	OWO WAT V 627	OWO WAT V 628	OWO WAT V 629	>	OWO WAT V 631	OWO WAT V 632	OWO WAT V 633	OWO WAT V 634	OWO WAT V 635	WAT V	OWO WAT V 637	OWO WAT V 638	OWO WAT V 639	OWO WAT V 640	OWO WAT V 641	WAT V	OWO WAT V 643	WAT V	WAT V	WAT V	WAT V	WAT V	WAT V	WAT V		WAT V	WAT V	OWO WAT V 654	OWO WAT V 655	WAT V	OWO WAT V 657	-	OW0 WAT V 659
0909	6061	6062	6063	6064	6065	9909	6067	6909	6909	6070	6071	6072	6073	6074	6075	9209	6077	6078	6019	6080	6081	6082	6083	6084	6085	9809	6087	6088	6083	0609	6091	6092	6093	6094	6095	9609	6097	8609	6609	6100	6101
œ ·	<b>&amp;</b>	8	œ	83	œ	æ	æ	œ	80	æ	89	œ	œ	æ	ထ	80	œ	œ	œ	œ	œ	æ	80	œ	œ	œ	α,	œ	œ	œ	œ	ω	00	œ	œ	æ	<b>&amp;</b>	œ	œ	ω	œ
1.00 35.51	1.00 34.53	1.00 35.10		1.00 34.54	1.00 35.82	1.00 34.54		1.00 34.23	1.00 34.65	1.00 35.48	1.00 34.69	1.00 35.07	1.00 35.68	1.00 35.69	1.00 35.74	1.00 35.64	1.00 35.51	1.00 35.66	1.00 36.16	1.00 35.14	1.00 35.66	1.00 36.54	1.00 35.34	1.00 35.66	1.00 36.08	1.00 35.59	1.00 35.60					1.00 36.33	1.00 35.23	1.00 35.08		1.00 37.22	1.00 35.78	1.00 36.07	1.00 36.81	1.00 35.61	1.00 36.55
31.457		59.956	26.130	71.942	73.743	10.945	10.060	18.773	35.620	36.602	20.660	50.572	11.023	36.911	26.394	25.261	54.266	67.743	44.339	53.643	54.942	52.116	37.685	22.637	48.242	15.641	15.117	1.142	9.308	75.539	32.749	14.031	36.864	29.088	35.706	13.735	50.997	12.139	38.927	46.301	64.153
61.496	53.881	68.150	41.546	74.712	73.783	55.518	60.566	63.329	54.481	63.600	96.750	73.376	50.506	50.549	55.539	93.666	73.996	48.697	101.046	69.586	79.574	80.128	51.551	70.631	48.081	81.332	74.805	73.953	85.286	52.963	66.230	44.107	86.958	69.199	50.939	58.135	55.369	52.384	103.615	79.803	67.375
0.045	46.395	52.136	19.009	34.720	37.259	27.833	46.309	10.952	-1.056	-4.178	4.690	8.484	44.017	3.744	49.672	27.932	22.451	33.048	12.820	51,198	29.662	32.247	6.142	5.622	22.365	8.288	10.941	35.796	38.777	20.560	-3.377	40.087	-1.635	48.364	18.383	18.334	16.341	28.580	23.108	12.325	21.280
OWO WAT V	OWO WAT V	OMO	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OMO	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT V	OWO WAT	OWO WAT V
6018	6019	6020	6021	6022	6023	6024	6025	6026	6027	6028	6029	6030	6031	6032	6033	6034	6035	6036	6037	6038	6039	6040	6041	6042	6043	6044	6045	6046	6047	6048	6049	6050	6051	6052	6053	6054	6055	9509	6057	6058	605

117 442.02 441.59 441.59 441.59 441.67 441.67 441.67 441.67 441.67 441.67 442.68 442.68 442.69 442.69 442.69 443.69 443.19 443.19 4.4 23.558 118.755 56.124 50.393 24.221 24.221 49.471 16.222 113.933 67.808 86.053 13.933 67.808 18.650 9.706 18.650 1 61.813 17.363 61.642 1.386 37.683 76.751 19.316 57.571 47.268 28.040 35.569 44.845 88.109 57.071 62.124 108.235 71.200 51.726 55.447 79.694 84.285 81.759 67.815 74.685 67.815 74.685 67.815 74.685 7 19.476
46.461.294
8.461.294
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8.481
8. 5228 5228 5229 5229 5229 5230 18.950 73.677 10.040 11.065 11.065 11.065 11.065 11.065 11.065 11.065 11.065 11.065 11.065 11.065 11.065 11.065 12.009 13.51 1 85.408
66.169
83.519
67.552
72.699
65.503
67.341
67.341
67.341
67.341
67.341
67.341
67.341
67.341
67.341
67.341
67.341
67.341
67.341
67.341
67.341
67.341
67.341
67.341
67.341 39.451 3.185 15.866 42.027 41.063 15.728 -8.665 22.964 52.964 52.964 52.964 62.001 4 8116 27.033 11.368 50.379 9.739 26.770 51.219 44.665 5.325 41.721 16.993 -3.245 10.854 6.873 6.873 10.333 25.342 11.886 20.669 12.363 13.886 20.669 13.389 25.342 13.389 25.342 13.389 27.369 20.669 20.669 20.699 21.098 

118 49.410 16.034 25.0411 49.410 49.520 49.520 49.520 49.520 66.580 66.380 33.730 33.730 33.730 33.730 43.685 54.685 5 45.863
47.238
90.618
73.271
50.245
64.072
64.107
74.829
96.027
94.673
100.590
106.460
93.590
66.093
97.178
93.590
66.093
93.590
72.473
49.673
72.566
56.933
49.673
72.473
49.673
72.473
49.673
72.473
49.673
72.473
49.673
72.566
89.920
71.480
73.136
89.154 4.605 42.143 26.105 3.202 -2.318 27.101 14.712 46.781 -6.114 47.645 48.802 6.382 20.784 15.678 43.426 10.651 0.918 24.371 7.425 20.314 12.244 2.847 15.592 45.552 48.867 51.123 28.449 31.691 45.269 118.096 -5.062 117.006 39.580 116.405 28.253 26.165 25.024 > > > ИАТ ИИАТ ИИОТ ИИОТ ИИОТ ИООТ 6342 6344 6345 6346 6346 6348 6349 6350 6351 43.38 44.1184 44.1184 44.1173 44.1184 44.11 000.11.000 43.918 11.170 14.779 16.127 34.245 78.143, 51.513 47.505 47.505 47.505 47.327 24.292 24.292 24.292 24.292 25.148 34.805 34.805 34.805 55.531 61.344 6 19.292 19.941 46.683 26.955 52.657 4.982 16.121 68.046 63.513 97.468 69.989 50.802 102.118 52.205 61.154 71.909 56.649 48.499 53.767 85.158 57.833 78.070 53.418 82.502 62.765 80.451 69.957 69.957 53.593 85.638 85.638 85.638 87.015 81.361 97.353 83.741 89.452 48.418 65.082 70.931 49.445 64.035 94.650 52.807 49.582 61.611 21.480 14.604 14.875 3.701 17.294 17.295 17. 51.902 14.161 0.976 11.378 23.725 31.619 21.564 

119 55.58 56.11 56.11 57.48 58.08 58.08 58.08 58.09 58 14.354 48.028 23.242 56.998 12.082 17.487 7.934 44.072 44.072 44.072 19.393 32.815 33.815 33.815 33.815 34.072 34.072 34.072 34.072 37.082 37. 48.219
49.394
105.611
51.479
51.479
61.879
46.684
46.516
92.028
89.221
108.504
94.059
93.333
94.684
94.253
93.343
94.253
94.253
94.253
94.253
94.253
94.253
94.253
94.253
94.253
94.253
94.253
94.253
94.253
94.253 49.464
-2.822.263
823.222.263
823.222.263
823.222.263
823.222.263
823.222.263
823.222.263
823.223.263
823.223.263
823.223.263
823.223.263
823.223.263
823.223.263
823.223.263
823.223.263
823.223.263
823.223.263
823.223.263
823.223.263
823.223
823.223
823.223
823.223
823.223
823.223
823.223
823.223
823.223
823.223
823.233
823.233  $\begin{smallmatrix} 7 & 1 & 1 & 1 & 1 & 1 \\ 2 & 1 & 2 & 1 & 1 & 1 \\ 2 & 1 & 2 & 1 & 1 \\ 2 & 1 & 2 & 1 & 1 \\ 2 & 1 & 2 & 1 & 1 \\ 2 & 1 & 2 & 1$ 53396 53997 5400 48.25 43.39 49.78 46.50 46.50 48.63 69.93 22.583
62.131
63.179
38.793
38.793
38.793
39.793
31.948
81.5810
16.693
11.916
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529
19.529 50.152
57.487
66.185
67.487
74.287
74.287
74.287
74.923
104.557
79.410
90.457
72.576
59.810
98.214
88.334
62.936
64.779
98.214
88.534
62.936
64.779
96.936
76.486
91.942
76.486
91.942
76.486
91.942
76.938
83.1033
83.1033
83.1033
83.1033
84.962
95.134
96.956
76.486
91.942
76.956
91.942 49.209
10.266
110.266
119.083
139.385
110.083
139.385
110.259
110.259
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289
120.289 5.3.54 5.3.54 5.3.54 5.3.55 5.3.55 5.3.55 5.3.55 5.3.55 5.3.55 6.3.55

120 56.21 3.99 8.928 43.406 42.398 30.747 -0.279 34.862 34.725 33.234 62.190 76.111 31.368 28.522 34.298 65.629 65.629 37.673 39.760 19.244 30.750 19.244 30.750 19.244 30.750 19.244 30.750 19.244 30.750 19.244 30.750 19.244 30.750 19.244 30.750 19.244 30.750 19.244 30.750 21.335 22.20 21.335 22.20 23.2 48.935
51.440
65.520
65.520
65.520
838.131
883.783
883.783
89.732
62.293
90.751
90.751
90.733
90.730
92.730
92.730
92.730
92.730
92.730
92.730
92.730
92.730
93.952
93.952
93.952
93.952
93.952
93.952
93.952
93.952
93.952
93.952 38.033 22.720 16.160 22.720 30.609 30.609 30.609 31.440 35.805 37.344 35.805 37.344 35.805 37.344 35.805 37.344 35.805 37.344 36.909 37.344 37.346 64880 64881 64885 64886 64886 64886 64896 6496 64996 6496 64996 6496 64996 6496 6496 6496 6496 6496 6496 649 60.39 61.97 47.55 88.13 66.13 47.55 86.44 66.13 66.44 1.1.000 1.1 12.475 68.228 84.75 12.286 13.27 13.27 13.27 13.27 13.27 14.11 14.11 14.11 14.11 15.19 16.19 17.60 17.60 17.60 18.31 17.60 17.60 18.31 18.31 18.71 18. 112. 668. 34. 34. 32. 35. 256. 228. 52.799
50.122
60.942
99.863
94.100
68.477
80.622
47.019
70.552
61.460
42.378
42.378
42.378
44.358
102.526
86.155
99.271
68.479
88.126
62.726
90.680
65.150
66.1182
90.680 24.857 24.857 26.545 26.545 28.187 41.335 29.677 10.580 29.677 29.677 20.029 39.076 41.636 44.085 26.670 24.479 24.479 24.479 24.479 25.549 39.076 39.076 39.076 39.076 39.076 39.076 39.077 39.076 39.077 5.4238 5.44338 5.44433 5.44433 6.4443 6.4 WO 99/43794

PCT/DK99/00088

121

 6522
 ONO WAT W 180
 51.301
 65.512
 30.778
 1.00 58.87

 6523
 ONO WAT W 181
 49.729
 67.077
 23.232
 1.00 56.44

 6524
 ONO WAT W 181
 15.428
 67.077
 23.232
 1.00 50.64

 6525
 ONO WAT W 181
 28.3127
 67.077
 21.323
 1.00 50.64

 6526
 ONO WAT W 183
 28.3127
 65.725
 1.00 52.88

 6527
 ONO WAT W 186
 39.667
 45.777
 21.789
 1.00 52.88

 6528
 ONO WAT W 186
 39.667
 45.777
 21.789
 1.00 52.28

 6523
 ONO WAT W 186
 49.525
 52.21
 11.484
 1.00 52.18

 6531
 ONO WAT W 189
 49.525
 52.21
 11.484
 1.00 52.18

 6532
 ONO WAT W 189
 49.525
 52.21
 11.484
 1.00 52.18

 6533
 ONO WAT W 190
 19.774
 47.99
 10.00 54.90
 10.00 54.90

 6534
 ONO WAT W 190
 19.774
 47.879
 10.00 54.90
 10.00 54.90

## **CLAIMS:**

5

15

25

30

- 1. A method of producing a variant of a parent maltogenic alpha-amylase, said method comprising
  - a) modeling the parent alpha-amylase on the three-dimensional structure of SEQ ID NO: 1 depicted in the Appendix to produce a three-dimensional structure of the parent alpha-amylase;
    - b) identifying in the three-dimensional structure obtained in step (a) at least one structural part of the parent wherein an alteration in said structural part is predicted to result in said altered property;
- c) modifying the sequence of a nucleic acid encoding the parent alphaamylase to produce a nucleic acid encoding a deletion, insertion, or substitution of one or more amino acids at a position corresponding to said structural part; and
  - d) expressing the modified nucleic acid in a host cell to produce the variant alpha-amylase,

wherein the variant has alpha-amylase enzymatic activity and has at least one altered property relative to the parent.

- The method of claim 1, wherein the altered property is pH dependent activity, thermostability, substrate cleavage pattern, specific activity of cleavage, transglyco-sylation, ability to reduce retrogradation of starch, ability to reduce staling of bread, substrate specificity, substrate binding or calcium binding.
  - 3. A method of constructing a variant of a parent maltogenic alpha-amylase, which method comprises:
    - a) identifying an amino acid residue which is within 15 Å (in particular 10 Å)
      from an active site residue of the parent amylase in the three-dimensional
      structure of said parent, and which is involved in electrostatic or hydrophobic
      interactions with an active site residue;
      - b) substituting said amino acid residue with another amino acid residue which changes the electrostatic and/or hydrophobic surroundings of an active site residue, and which can be accommodated in the structure;
      - c) optionally repeating steps a) and b) recursively;
      - d) optionally, making alterations each of which is an insertion, a deletion or a substitution of an amino acid residue at one or mor positions other than b),

5

10

15

20

30

- e) preparing the variant resulting from steps a) d);
- f) testing the pH dependent activity of said variant; and
- g) optionally repeating steps a) f) recursively; and
- h) selecting a variant having an altered pH dependent activity as compared to the parent amylase.
- 4. A method of constructing a variant of a parent maltogenic alpha-amylase, which method comprises:
  - a) identifying an internal cavity or crevice in the three-dimensional structure of said parent;
  - b) substituting an amino acid residue in the neighborhood of the cavity or crevice with another amino acid residue which increases the hydrophobic interaction and/or fills out or reduces the size of the cavity or crevice;
    - c) optionally repeating steps a) and b) recursively;
- d) optionally, making alterations each of which is an insertion, a deletion or a substitution of an amino acid residue at one or more positions other than b),
  - e) preparing the variant resulting from steps a) d);
  - f) testing the thermostability of said variant; and
  - g) optionally repeating steps a) f) recursively; and
  - h) selecting a variant having increased thermostability as compared to the parent amylase.
- 5. The method of claim 4, wherein the substitution of the amino acid residue results in increasing the hydrophobic interaction, substitution with proline, substitution of histidine with another amino acid, stabilization of calcium binding, introduction of an interdomain disulfide bond, removal of a deamidation site, altering a hydrogen bond contact, filling in an internal structural cavity with an amino acid with a bulkier side group, introduction of interdomain interactions, altering charge distribution, helix capping, or introduction of a salt bridge.
  - 6. A method of constructing a variant of a parent maltogenic alpha-amylase, which method comprises:
    - a) identifying an amino acid residue within 10 Å from a calcium binding site in the three dimensional structure of the amylase;
    - b) substituting the amino acid residue with another amino acid residue so as to improve the interaction with the calcium ion;
    - c) optionally rep ating steps a) and b) recursively;

5

- d) optionally, making alterations each of which is an insertion, a deletion or a substitution of an amino acid residue at one or more positions other than b),
- e) preparing the variant resulting from steps a) d);
- f) testing the thermostability of said variant; and
- g) optionally repeating steps a) f) recursively; and
  - h) selecting a variant having increased thermostability as compared to the parent amylase.
- 7. A method of constructing a variant of a parent maltogenic alpha-amylase, which method comprises:
- a) identifying the substrate binding area in a model of the three-dimensional structure of the parent amylase;
  - b) modifying the substrate binding area by an amino acid substitution, deletion or insertion;
  - c) optionally repeating step b) recursively;
- d) optionally, making alterations each of which is an insertion, a deletion or a substitution of an amino acid residue at one or more positions other than b),
  - e) preparing the variant resulting from steps a) d);
  - f) testing the substrate-cleavage pattern of the variant.
  - g) optionally repeating steps a) f) recursively; and
- 20 h) selecting a variant having an altered substrate-cleavage pattern as compared to the parent amylase.
  - 8. A method for producing a maltogenic alpha-amylase variant, comprising:
    - a) constructing the variant by the method of any of claims 2-7;
    - b) transforming a microorganism with a DNA sequence encoding the variant;
- c) cultivating the transformed microorganism under conditions which are conducive for producing the variant, and
  - d) optionally, recovering the variant from the resulting culture broth.
  - 9. A polypeptide which:
    - a) has maltogenic alpha-amylase activity;
- b) has at least 70 % identity to SEQ ID NO: 1;
  - c) comprises an amino acid modification compared to SEQ ID NO: 1 at a position corresponding to D127, V129, F188, A229, Y258, V281, F284, T288, N327, M330, G370, N371, and/or D372; and
- d) has altered pH dependent activity as compared to the polypeptide of SEQ
   ID NO: 1.

- 10. The polypeptide of claim 9, wherein the modification comprises a substitution corresponding to D127N/L, V129S/T/G/V, F188E/K/H, A229S/T/G/V, Y258E/D/K/R/F/N, V281L/T, F284K/H/D/E/Y, T288E/K/R, N327D, M330L/F/I/D/E/K, G370N, N371D/E/G/K, and/or D372N/V.
- 5 11. A polypeptide which:
  - a) has maltogenic alpha-amylase activity;
  - b) has at least 70 % identity to SEQ ID NO: 1; and
- c) comprises an amino acid modification compared to SEQ ID NO: 1 at a position corresponding to Q13, I16, D17, N26, N28, P29, A30, S32, Y33, G34, L35, K40, M45, P73, V74, D76, N77, D79, N86, R95, N99, I100, H103, Q119, N120, N131, S141, T142, A148, N152, A163, H169, N171, G172, I174, N176, N187, F188, A192, Q201, N203, H220, N234, G236, Q247, K249, D261, N266, L268, R272, N275, N276, V279, N280, V281, D285, N287, F297, Q299, N305, K316, N320, L321, N327, A341, N342, A348, Q365, N371, N375, M378, G397, A381, F389, N401, A403, K425, N436, S442, N454, N468, N474, S479, A483, A486, V487, S493, T494, S495, A496, S497, A498, Q500, N507, I510, N513, K520, Q526, A555, A564, S573, N575, Q581, S583, F586, K589, N595, G618, N621, Q624, A629, F636, K645, N664 and/or T681; and
  - d) has improved stability compared to the polypeptide of SEQ ID NO: 1.
- The polypeptide of claim 11, wherein the modification comprises at a position corresponding to K40, V74, H103, S141, T142, F188, H220, N234, K249, D261, L268, V279, N342, H344, G397, A403, K425, S442, S479, S493, T494, S495, A496, S497, A498, Q500, K520, A555 and/or N595; preferably a substitution corresponding to K40R, V74P, H103Y/V/I/L/F/Y, S141P, T142A, F188I/L, H220Y/L/M, N234P, K249P, D261G, L268P, V279P, N342P, H344E/Q/N/D/Y, G397P, A403P, K425E, S442P, S479P, S493P, T494P, S495P, A496P, S497P, A498P, Q500P, K520R, A555P and/or N595I.
- 13. The polypeptide of claim 11 or 12, wherein the modification comprises at a position corresponding to D17, N28, P29, A30, S32, Y33, G34, R95, H103, N131, 30 H169, I174 and/or Q201 such as to improve calcium coordination, preferably a substitution corresponding to D17Q/E, A30D/M/L/AV/I/E/Q, S32D/E/N/Q, R95M/L/AV/I/E/Q, H103Y/N/Q/D/E, N131D, H169N/D/E/Q, I174E/Q, Q201E.
  - 14. The polypeptide of any of claims 11-13, wherein the modification comprises a substitution at a position corresponding to Q13, N26, N77, N86, N99, Q119, N120,

N131, N152, N171, N176, N187, Q201, N203, N234, Q247, N266, N275, N276, N280, N287, Q299, N320, N327, N342, Q365, N371, N375, N401, N436, N454, N468, N474, Q500, N507, N513, Q526, N575, Q581, N621, Q624 and/or N664 such as to remove a deamidation site, preferably a substitution corresponding to 5 Q13S/T/AV/L/I/F/M, N26S/T/AV/L/I, N77S/T/AV/L/I, N86S/T/AV/L/I, N99T/S/V/L, N120S/T/A/V/L/I, N131S/T/A/V/L/I, N152T/S/V/L, N171Y/D/S/T, Q119T/S, N187S/T/A/V/L/I, Q201S/T/A/V/L/I/F/M, N203D/S/T/AV/L/I. N176S/T/A/V/L/I. Q247S/T/A/V/L/I/F/M, N266S/T/A/V/L/I, N234S/T/A/V/L/I, N275S/T/A/V/L/I, N276S/T/A/V/L/I, N280S/T/A/V/L/I, N287S/T/A/V/L/I, Q299L/T/S, N320S/T/A/V/L/I, N342S/T/A/V/L/I, Q365S/T/A/V/L/I. N371S/T/A/V/L/I. 10 N327S/T/AV/L/I. N401S/T/A/V/L/I, N436S/T/A/V/L/I, N454D/S/T/A/V/L/I, N375S/T/A/V/L/I, N474D/S/T/A/V/L/I, Q500S/T/A/V/L/I/F/M, N507S/T/A/V/L/I, N468D/S/T/A/V/L/I. N513S/T/A/V/L/I, Q526 D/S/T/A/V/L/I, N575S/T/A/V/L/I, Q581S/T/A/V/L/I/F/M, N621S/T/A/V/L/I Q624S/T/A/V/L/I/F/M and/or N664D/S/T/A/V/L/I.

- The polypeptide of any of claims 11-14, wherein the modification comprises a substitution at a position corresponding to I16, L35, M45, P73, D76, D79, A192, I100, A148, A163+G172, L268, V281, D285, L321, F297, N305, K316, S573, A341, M378, A381, F389, A483, A486, I510, A564, F586, K589, F636, K645, A629, and/or T681 such as to improve hydrogen bond contact, preferably a substitution corresponding to I16T/D/N, L35Q, M45K, P73Q, D76E, D79E/Y, A192S/D/N, I100T/S/D/N/E/Q, A148D/N/E/Q/S/T/R/K, A163Y+G172S/D/N, L268R/K, V281/Q, D285R/K, L321Q, F297N/D/Q/E, N305K/R, K316N/D, S573N/D, A341R/K, M378R/K, A381S/D/N, F389Y, A483S/D/N, A486Q/E, I510R/K, A564S/D/N, F586S/D/N, 'K589S/D/Q/N, F636Y, K645T, A629N/D/E/Q, and/or T681D/N/E/Q/S.
- 25 16. The polypeptide of any of claims 11-15, wherein the modification comprises substitutions such as to introduce one or more interdomain disulfide bonds, preferably corresponding to G236C + S583C, G618C + R272C, and/or A348C + V487C.
- The polypeptide of any of claims 11-16, wherein the substitution at a position corresponding to L51, L75, L78, G88, G91, T94, V114, I125, V126, T134, G157, L217, S235, G236, V254, V279, V281, L286, V289, I290, V308, L321, I325, D326, L343, F349, S353, I359, I405, L448, Q449, L452, I470, G509, V515, S583, G625, L627, L628 and/or A670 so as to fill an internal cavity or crevice, preferably a substitution corresponding to L51W, L75F/Y, L78I, G88A/V/T, G91T/S/V/N, T94V/I/L, V114V/I/L, I125L/M/F/Y/W, V126I/L, T134V/I/L/M/F/Y/W, G157A/V/I/L,
- 35 L217V/I/M/F/Y/W, S235I/L/M/F/Y/W, G236A/V/I/L/M/F/Y/W, V254I/L/M/F/Y/W,

V279M/I/L/F, V281I/L/M/F/Y/W, L286F, V289I/L/R, I290M/L/F, V308I/L/M/F/Y/W, L321I/M/F/Y/W, I325L/M/F/Y/W, D326E/Q, L343M/F/Y/W, F349W/Y, S353V/I/L, I359L/M/F/Y/W, I405M/L/Y/F/W, L448Y, Q449Y, L452M/Y/F/W, I470M/L/F, G509A/V/I/L/M/S/T/D/N, V515I/L, S583V/I/L/V, G625A/V/I/L/M/F/Y/W, L627M/F/Y, L628M/I/F/Y/W, A670V/I/L/M/F/Y/W and/or L217 in combination with L75 (e.g. L217F/Y in combination with L75F/Y).

- 18. The polypeptide of any of claims 11-17, wherein the modification comprises a substitution at a position corresponding to N106, N320 and Q624 so as to create a salt bridge, preferably a substitution corresponding to N106R, N320E/D and/or Q624E.
  - 19. The polypeptide of any of claims 11-18, wherein the modification comprises a substitution at a position corresponding to K244 and/or K316 such as to alter the charge distribution, preferably a substitution corresponding to K244S and/or K316G/N/D.
- 15 20. The polypeptide of any of claims 11-19, wherein the modification comprises a substitution at a position corresponding to V281 and/or A629 such as to alter the binding site, preferably a substitution corresponding to V281Q and/or A629N/D/E/Q.
- 21. The polypeptide of any of claims 11-20, wherein the modification comprises substitutions such as to alter the interdomain interaction at a position corresponding to F143+F194+L78, A341+A348+L398+l415+T439+L464+L465, L557, S240+L268, Q208+L628, F427+Q500+N507+M508+S573 and/or I510+V620, preferably substitutions corresponding to F143Y+F194Y+L78Y/F/W/E/Q, A341S/D/N+A348V/I/L+L398E/Q/N/D+I415E/Q+T439D/E/Q/N+L464D/E+L465D/E/N/Q/R/K, L557Q/E/N/D, S240D/E/N/Q+L268D/E/N/Q/R/K, Q208D/E/Q+L628E/Q/N/D, F427E/Q/R/K/Y+Q500Y+N507Q/E/D+M508K/R/E/Q+S573D/E/N/Q; and/or I510D/E/N/Q/S+V620D/E/N/Q.
  - 22. A polypeptide which:
    - a) has maltogenic alpha-amylase activity;
    - b) has at least 70 % identity to SEQ ID NO: 1;
- c) comprises an amino acid modification compared to SEQ ID NO: 1 at a position corresponding to P191, A192, G193, F194 and/or S195; and
  - d) has higher specific amylase activity than the polypeptide of SEQ ID NO:
  - 1.

- 23. The polypeptide of claim 22, wherein the modification comprises a deletion, preferably the deletion  $\Delta$  (191-195).
- 24. The polypeptide of claim 22, wherein the modification comprises insertion, preferably 192-A-193.
- 5 25. A polypeptide which:

10

25

- a) has maltogenic alpha-amylase activity;
- b) has at least 70 % identity to SEQ ID NO: 1;
- c) comprises an amino acid modification compared to SEQ ID NO: 1 at a position corresponding to A30, K40, N115, T142, F188, T189, P191, A192, G193, F194, S195, D261, T288, N327, K425, K520 and/or N595; and
- d) has a higher ability than the polypeptide of SEQ ID NO: 1 to reduce retrogradation of starch and/or staling of bread.
- The polypeptide of claim 26, wherein the modification comprises A30D, K40R, N115D, T142A F188L, T189Y, Δ (191-195), D261G, T288P, N327S, K425E, K520R
   and/or N595I.
  - 27. A nucleic acid sequence encoding the polypeptide of any of claims 9-26, preferably operably linked to one or more control sequences which direct the expression of the variant in a suitable expression host.
- 28. A recombinant expression vector comprising the nucleic acid sequence of claim 27, a promoter, and transcriptional and translational stop signals, and preferably further comprising a selectable marker.
  - 29. A transformed host cell comprising the nucleic acid sequence of claim 27 or the vector of claim 28.
  - 30. A method for producing the polypeptide of any of claims 9-26, comprising:
  - a) cultivating the transformed host cell of claim 29 under conditions conducive to expression of the variant; and
    - b) recovering the variant.
  - 31. A process for preparing a dough or a baked product prepared from the dough which comprises adding the polypeptide of any of claims 9-26, or a variant produced

WO 99/43794

PCT/DK99/00088

129

by the method of any of claims 1-8 to the dough in an amount which is effective to retard the staling of the bread.

32. The process of claim 31, wherein the variant is added in an amount of 0.1-5 mg per kg of flour, preferably 0.5-2 mg/kg.

1/1

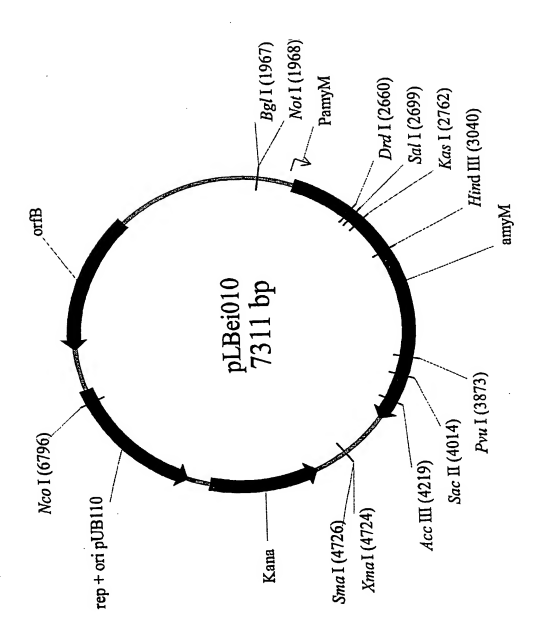


Fig. 1

WO 99/43794

PCT/DK99/00088

## SEQUENCE LISTING

<110> NOVO NORDISK A/S
<120> MALTOGENIC ALPHA-AMYLASE VARIANTS
<130> 5443-WO, SLK
<140>
<141>
<160> 14
<170> PatentIn Ver. 2.0
<210> 1
<211> 2160
<212> DNA
<213> Bacillus sp.
<220>
<221> CDS
<222> (1)(2160)
<220>
<221> mat_peptide
<222> (100)(2157)
<400> 1 atg as as as as ctt tct tta ttt gtg gga ctg atg ctc ctc atc 48
Met Lys Lys Lys Thr Leu Ser Leu Phe Val Gly Leu Met Leu Leu Ile
-30 -25 -20
ggt ctt ctg ttc agc ggt tct ctt ccg tac aat cca aac gcc gct gaa 96
Gly Leu Leu Phe Ser Gly Ser Leu Pro Tyr Asn Pro Asn Ala Ala Glu
-15 -10 -5
gcc agc agt tcc gca agc gtc aaa ggg gac gtg att tac cag att atc 14
gcc agc agt tcc gca agc gtc aaa ggg gac gtg att tac cag att atc 14.  Ala Ser Ser Ser Ala Ser Val Lys Gly Asp Val Ile Tyr Gln Ile Ile
-1 1 5 10 15
att gac egg ttt tac gat ggg gac acg acg aac aac aat cet gec aaa 19
Ile Asp Arg Phe Tyr Asp Gly Asp Thr Thr Asn Asn Asn Pro Ala Lys
20 25 30
agt tat gga ctt tac gat ccg acc aaa tcg aag tgg aaa atg tat tgg 24 Ser Tyr Gly Leu Tyr Asp Pro Thr Lys Ser Lys Trp Lys Met Tyr Trp

ggc	ggg	gat	ctg	gag	ggg	gtt	cgt	caa	aaa	ctt	cct	tat	ctt	aaa	cag	288
Gly	Gly	Asp	Leu	Glu	Gly	Val	Arg	Gln	Lys	Leu	Pro	Tyr	Leu	Lys	Gln	
		50					55					60				
ctg	ggc	gta	acg	aca	atc	tgg	ttg	tcc	ccg	gtt	ttg	gac	aat	ctg	gat	336
Leu	Gly	Val	Thr	Thr	Ile	Trp	Leu	Ser	Pro	Val	Leu	Asp	Asn	Leu	Asp	
	65					70					75					
aca	ctg	gcg	ggc	acc	gat	aac	acg	ggc	tat	cac	gga	tac	tgg	acg	cgc	384
Thr	Leu	Ala	Gly	Thr	Asp	Asn	Thr	Gly	Tyr	His	Gly	Tyr	Trp	Thr	Arg	
80			_		85					90					95	
gat	ttt	aaa	cag	att	gag	gaa	cat	ttc	ggg	aat	tgg	acc	aca	ttt	gac	432
-										Asn						
		•		100					105		_			110		
acq	ttg	gtc	aat	gat	gct	cac	caa	aac	gga	atc	aag	gtg	att	gtc	gac	480
_	_	_		_	_					Ile	_		_		_	
			115	-				120			-		125		_	
ttt	ata	ccc	aat	cat	tcg	act	cct	ttt	aag	gca	aac	gat	tcc	acc	ttt	528
	-				_				_	Ala						
		130					135		•			140				
aca	gaa	ggc	gge	acc	ctc	tac	aac	aat	gga	acc	tat	ato	aac	aat	tat	576
	_			-						Thr			_			
	145	1	2			150					155		•			
ttt	gat	gac	aca	aca	aaa	aaa	tac	ttc	cac	cat	aat	aaa	gac	atc	agc	624
	_	_	_							His			-			
160					165	•	•			170			•		175	
										•						
ABC	taa	GAC	gac	caa	tac	σaσ	aca	CAA	taa	aaa	aac	ttc	aco	gat	cca	672
		-								Lys						
				180	-2-				185	_				190		
acc	aat	ttc	tca	ctt	acc	gat	tta	tca	cag	gaa	aat	ggc	aco	att	act	720
_								_	_	Glu					Ala	
			195					200				2	205			
caa	tac	cta	acc	gat	aca	aca	att	caa	tta	gta	qca	cat	qqa	aca	gat	768
		_		_			-		_	. Val	_				-	
	-3-	210					215					220	3			
aa+	tta	caa	att	gat	aca	ata	880	cat	ttt	ast	tea	aaa	tte	tee	aaa	816
										Asn						
1	225	_				230	_				235				-, -	

tcg	ttg	gcc	gat	aaa	ctg	tac	caa	aag	aaa	gac	att	ttc	ctg	gtg	ggg	864
Ser	Leu	Ala	Asp	Lys	Leu	Tyr	Gln	Lys	Lys	Asp	Ile	Phe	Leu	Val	Gly	
240					245					250					255	
gaa	tgg	tac	gga	gat	gac	ccc	gga	aca	gcc	aat	cat	ctg	gaa	aag	gtc	912
Glu	Trp	Tyr	Gly	Asp	Asp	Pro	Gly	Thr	Ala	Asn	His	Leu	Glu	Lys	Val	
	-	-	-	260	_		_		265					270		
cqq	tac	gcc	aac	aac	agc	ggt	gtc	aat	gtg	ctg	gat	ttt	gat	ctc	aac	960
Arg	Tvr	Ala	Asn	Asn	Ser	Glv	Val	Asn	Val	Leu	Asp	Phe	Asp	Leu	Asn	
	-4-		275					280					285			
acq	ata	att	cqa	aat	gtg	ttc	ggc	aca	ttt	acq	caa	acq	atq	tac	gat	1008
_			_							-	Gln		_		_	
		290	5				295					300		- , -		
												500				
ctt	aac	aat	ato	ata	aac	caa	aco	aaa	aac	gag	tac	222	tac	***	gaa	1056
			_								Tyr				_	2000
204	305	#P	2200	• • •		310		027	****	<b>514</b>	315	my G	-1-	-y s	914	
	303					320					243					
aat	cta	atc	aca	ttt	atc	gat	aac	cat	gat	atσ	tca	aga	ttt	ctt	tea	1104
						-			-	_	Ser	_			-	
320				• • • • •	325					330		9		~~~	335	
320										330					333	
σta	aat	tca	aac	аас	aca	aat	tta	cac	cag	aca	ctt	get	ttc	att	ete	1152
_		_		_			_		_		Leu	-				
				340					345					350		
				340					343					330		
act	tcq	caa	ggt	acg	ccc	tcc	atc	tat	tat	qqa	acc	qaa	caa	tac	ato	1200
	_										Thr	_			_	
			355					360	•				365			
qca	qqc	ggc	aat	gac	ccg	tac	aac	cgq	ggg	atq	atg	CCQ	aca	ttt	gat	1248
-				_	_					_	Met	_				
		370		•		•	375					380				
		• • •														
acq	aca	acc	acc	acc	ttt	aaa	gag	ata	tca	act	ctg	aca	aaa	tta	cac	1296
								_			Leu			_	_	
	385					390					395		017		,,,,	
	505					330					333					
agg	aac	aat	gca	gca	atc	cao	tac	dac	acc	acc	acc	cao	cat	taa	atc	1344
											Thr					
400					405		-1-	1		410		7111	9	-+1	415	
aac	aat	gat	att	tac	att	tat	GAR	Caa	888	tee	tte	880	get	atc	gtg	1392
											Phe					
				420		- ,		5	425		- 444		بإد	430		

ttg	gtg	gcc	atc	aat	cga	aac	acg	caa	tcc	tcc	tat	tcg	att	tcc	ggt	1440
Leu	Val	Ala	Ile	Asn	Arg	Asn	Thr	Gln	Ser	Ser	Tyr	Ser	Ile	Ser	Gly	
			435					440					445			
tta	cag	aco	qec	tta	cca	aat	aac	agc	tat	aca	gat	tat	cta	tca	aaa	1488
			Ala	_				_			•		_			
200		450					455		-1-			460			2	
		430					133									
														-a-		1536
_	_		ggg					_							_	1336
Leu		GIĀ	Gly	ABN	GIY		ser	VAI	Ser	ABN	_	ser	Val	AIA	ser	
	465					470					475					
	_		gcg			_				-						1584
Phe	Thr	Leu	Ala	Pro	Gly	Ala	Val	Ser	Val	Trp	Gln	Tyr	Ser	Thr	Ser	
480					485					490					495	
gct	tca	gcg	ccg	caa	atc	gga	tcg	gtt	gct	cca	aat	atg	ggg	att	ccg	1632
Ala	Ser	Ala	Pro	Gln	Ile	Gly	Ser	Val	Ala	Pro	Asn	Met	Gly	Ile	Pro	
				500					505					510		
aat	aat	ata	gtc	acq	atc	gac	aaa	aaa	aat	ttt	aaa	acq	acq	caq	gga	1680
			Val	_		_						_	_	_		
,			515					520	,		3		525		3	
		200	ttt	~~~	~~~	~-~		~~~	205	~+~		***	+~~	200	tac	1728
																1/20
Thr	val		Phe	GIY	GIY	AST		Ala	Inr	VAI	ьўв		11p	Int	Ser	
		530					535					540				
			gaa	-		_			_	_	_		_		_	1776
Asn	Arg	Ile	Glu	Val	Tyr	Val	Pro	Asn	Met	Ala	Ala	Gly	Leu	Thr	ДаД	
	545					550					555					
gtg	aaa	gtc	acc	gcg	ggt	gga	gtt	tcc	agc	aat	ctg	tat	tet	tac	aat	1824
Val	Lys	Val	Thr	Ala	Gly	Gly	Val	Ser	Ser	Asn	Leu	Tyr	Ser	Tyr	Asn	
560					565					570					575	
att	ttg	agt	gga	acg	cag	aca	tcg	gtt	gtg	ttt	act	gtg	aaa	agt	gcg	1872
Ile	Leu	Ser	Gly	Thr	Gln	Thr	Ser	Val	Val	Phe	Thr	Val	Lys	Ser	Ala	
			_	580					585				_	590		
cat	cca	acc	aac	cta	gae	gat	aac	att	tac	cto	acc	ggc	aac	ata	ccg	1920
	_		Asn	_		-	_			_	_				_	
210	-10	- 111	595	200	CIY	woh	-y 5	600	_	<b>20</b> €	- 114	411	605	0		
			233					900					003			
																1060
_	_				_	-	_	_	_		-	_			gcg	1968
Glu	Leu	_	Asn	Trp	ser	Thr	_		ser	GIĀ	Ala		Asn	Asn	ATA	
		610					615					620				

cas ggg ccc ctg ctc gcg ccc sat tat ccg gat tgg ttt tat gta ttc Gln Gly Pro Leu Leu Ala Pro Asn Tyr Pro Asp Trp Phe Tyr Val Phe 625 630 age gtt cca gca gga aag acg att caa tte aag tte tte ate aag egt Ser Val Pro Ala Gly Lys Thr Ile Gln Phe Lys Phe Phe Ile Lys Arg 645 gcg gat gga acg att caa tgg gag aat ggt tcg aac cac gtg gcc aca Ala Asp Gly Thr Ile Gln Trp Glu Asn Gly Ser Asn His Val Ala Thr act ccc acg ggt gca acc ggt aac att act gtt acg tgg caa aac tag 2160 Thr Pro Thr Gly Ala Thr Gly Asn Ile Thr Val Thr Trp Gln Asn 680 <210> 2 <211> 719 <212> PRT <213> Bacillus sp. Met Lys Lys Thr Leu Ser Leu Phe Val Gly Leu Met Leu Leu Ile Gly Leu Leu Phe Ser Gly Ser Leu Pro Tyr Asn Pro Asn Ala Ala Glu 25 Ala Ser Ser Ser Ala Ser Val Lys Gly Asp Val Ile Tyr Gln Ile Ile Ile Asp Arg Phe Tyr Asp Gly Asp Thr Thr Asn Asn Asn Pro Ala Lys 55 Ser Tyr Gly Leu Tyr Asp Pro Thr Lys Ser Lys Trp Lys Met Tyr Trp Gly Gly Asp Leu Glu Gly Val Arg Gln Lys Leu Pro Tyr Leu Lys Gln Leu Gly Val Thr Thr Ile Trp Leu Ser Pro Val Leu Asp Asn Leu Asp 100 Thr Leu Ala Gly Thr Asp Asn Thr Gly Tyr His Gly Tyr Trp Thr Arg

120

Asp Phe Lys Gln Ile Glu Glu His Phe Gly Asn Trp Thr Thr Phe Asp Thr Leu Val Asn Asp Ala His Gln Asn Gly Ile Lys Val Ile Val Asp 150 Phe Val Pro Asn His Ser Thr Pro Phe Lys Ala Asn Asp Ser Thr Phe 170 Ala Glu Gly Gly Ala Leu Tyr Asn Asn Gly Thr Tyr Met Gly Asn Tyr Phe Asp Asp Ala Thr Lys Gly Tyr Phe His His Asn Gly Asp Ile Ser 200 Asn Trp Asp Asp Arg Tyr Glu Ala Gln Trp Lys Asn Phe Thr Asp Pro 215 Ala Gly Phe Ser Leu Ala Asp Leu Ser Gln Glu Asn Gly Thr Ile Ala 225 230 235 240 Gin Tyr Leu Thr Asp Ala Ala Val Gin Leu Val Ala His Gly Ala Asp 250 Gly Leu Arg Ile Asp Ala Val Lys His Phe Asn Ser Gly Phe Ser Lys 265 Ser Leu Ala Asp Lys Leu Tyr Gln Lys Lys Asp Ile Phe Leu Val Gly 280 Glu Trp Tyr Gly Asp Asp Pro Gly Thr Ala Asn His Leu Glu Lys Val 295 Arg Tyr Ala Asn Asn Ser Gly Val Asn Val Leu Asp Phe Asp Leu Asn 305 315 Thr Val Ile Arg Asn Val Phe Gly Thr Phe Thr Gln Thr Met Tyr Asp 325 330 Leu Asn Asn Met Val Asn Gln Thr Gly Asn Glu Tyr Lys Tyr Lys Glu 345 Asn Leu Ile Thr Phe Ile Asp Asn His Asp Met Ser Arg Phe Leu Ser 355 360 365 Val Asn Ser Asn Lys Ala Asn Leu His Gln Ala Leu Ala Phe Ile Leu

370

Thr Ser Arg Gly Thr Pro Ser Ile Tyr Tyr Gly Thr Glu Gln Tyr Met 385 390 395 400

- Ala Gly Gly Asn Asp Pro Tyr Asn Arg Gly Met Met Pro Ala Phe Asp 405 410 415
- Thr Thr Thr Ala Phe Lys Glu Val Ser Thr Leu Ala Gly Leu Arg
  420 425 430
- Arg Asn Asn Ala Ala Ile Gln Tyr Gly Thr Thr Thr Gln Arg Trp Ile 435 440 445
- Asn Asn Asp Val Tyr Ile Tyr Glu Arg Lys Phe Phe Asn Asp Val Val 450 455 460
- Leu Val Ala Ile Asn Arg Asn Thr Gln Ser Ser Tyr Ser Ile Ser Gly 465 470 475 480
- Leu Gln Thr Ala Leu Pro Asn Gly Ser Tyr Ala Asp Tyr Leu Ser Gly
  485 490 495
- Leu Leu Gly Gly Asn Gly Ile Ser Val Ser Asn Gly Ser Val Ala Ser 500 505 505
- Phe Thr Leu Ala Pro Gly Ala Val Ser Val Trp Gln Tyr Ser Thr Ser 515 520 525
- Ala Ser Ala Pro Gln Ile Gly Ser Val Ala Pro Asn Met Gly Ile Pro 530 535 540
- Gly Asn Val Val Thr Ile Asp Gly Lys Gly Phe Gly Thr Thr Gln Gly 545 550 555 560
- Thr Val Thr Phe Gly Gly Val Thr Ala Thr Val Lys Ser Trp Thr Ser 565 570 575
- Asn Arg Ile Glu Val Tyr Val Pro Asn Met Ala Ala Gly Leu Thr Asp 580 585 590
- Val Lys Val Thr Ala Gly Gly Val Ser Ser Asn Leu Tyr Ser Tyr Asn 595 600 605
- Ile Leu Ser Gly Thr Gln Thr Ser Val Val Phe Thr Val Lys Ser Ala 610 615 620
- Pro Pro Thr Asn Leu Gly Asp Lys Ile Tyr Leu Thr Gly Asn Ile Pro 625 630 630 635 640

Glu	Leu	Gly	Asn	Trp 645	Ser	Thr	Asp	Thr	Ser 650	Gly	Ala	Val	Asn	Asn 655	Ala	
Gln	Gly	Pro	Leu 660	Leu	Ala	Pro	Asn	Tyr 665	Pro	Авр	Trp	Phe	Tyr 670	Val	Phe	
Ser	Val	Pro 675	Ala	Gly	Lys	Thr	Ile 680	Gln	Phe	Гув	Phe	Phe 685	Ile	Lys	Arg	
Ala	Asp 690	Gly	Thr	Ile	Gln	Trp 695	Glu	Asn	Gly	Ser	Asn 700	His	Val	Ala	Thr	
Thr 705	Pro	Thr	Gly	Ala	Thr 710	Gly	Asn	Ile	Thr	Val 715	Thr	Trp	Gln	Asn		
<21 <21 <21	0>	8 NA rtif			of Ar		icial	L Sec	lnevo	ce: 1	F 188	3H P1	rimer	e		
	00> 3 Latg		aac	caca	cgg 8	atcca	agcc	gg c	ttot	ege						38
<2: <2: <2:		38 DNA	fici	al S	eque:	nce										
<2 <2	20> 23>	Desc	ript	ion	of A	rtif	icia	l Se	drez	ce:	F188	E Pr	imer	•		
	00> aatg		aac	gaga	ıcgg	atco	agco	gg (	ettet	cgc					D	38
<2 <2		36 DNA		ial :	Seque	ence										
<: <:	220> 223>	Des	crip	tion	of 2	Arti	fici	al S	edne	nce:	F 2	84E	Prim	er		

WO 99/43794	PCT/DK99/00
<400> 5	
ggtgtcaatg tgctggatga agatctcaac acggtg	36
<210> 6	
<211> 36	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> Description of Artificial Sequence: F 284D Primer	
<400> 6	
ggtgtcaatg ttctagatga tgatctcaac acggtg	36
<210> 7	
<211> 36	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> Description of Artificial Sequence: F 284K Primer	
<400> 7	
ggtgtcaatg tgctggataa agatctcaac acggtg	36
<210> 8	
<211> 35	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> Description of Artificial Sequence: N 327D Primer	
<400> 8	
cacatttatc gatgatcatg atatgtcaag atttc	35
210. 0	
<210> 9 <211> 34	
<211> 34 <212> DNA	
<213> Artificial Sequence	
errerorar peddauca	
<220>	

PCT/DK99/00088

<223> Description of Artificial Sequence: T 288K Primer

WO 99/43794	PCT/DK99/00088
<400> 9	
cctamaacta gagttgttee actaggeett acac	34
standard adaptation academic acad	34
<210> 10	
<211> 34	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> Description of Artificial Sequence: T 288R Primer	
<400> 10	
cctamaacta gagttgtccc actaggcctt acac	34
<210> 11	
<211> 22	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> Description of Artificial Sequence: A 189 Primer	
<400> 11	
tgggcaatta ttttgatgac gc	22
<210> 12	
<211> 20	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> Description of Artificial Sequence: B 649 Primer	
<400> 12	
tccgctcgta tccgtgctcc	20
<210> 13	
<211> 20	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> Description of Artificial Sequence: A 82 Primer	

WO 99/43794	
<400> 13	20
ggggatctgg agggggttcg	20
<210> 14	
<211> 22	
<212> DNA	
<213> Artificial Sequence	
<220>	
<223> Description of Artificial Sequence: B 346 Primer	
<400> 14	22
httetpeteg tteecegttt gg	44

PCT/DK99/00088



## INTERNATIONAL SEARCH REPORT

International application No. PCT/DK 99/00088

	PUITUR	99/00088
A. CLASSIFICATION OF SUBJECT MATTER		
IPC6: C12N 9/28, A21D 8/04 According to International Patent Classification (IPC) or to both n	ational classification and IPC	
B. FIELDS SEARCHED		
Minimum documentation searched (classification system followed b	y classification symbols)	
IPC6: C12N		
Documentation searched other than minimum documentation to the	e extent that such documents are in	cluded in the fields searched
SE,DK,FI,NO classes as above		
Electronic data base consulted during the international search (name	e of data base and, where practicable	le, search terms used)
C. DOCUMENTS CONSIDERED TO BE RELEVANT		
Category* Citation of document, with indication, where ap	propriate, of the relevant passag	ges Relevant to claim No.
X WO 9623874 A1 (NOVO NORDISK A/S (08.08.96), see page 30-37,	), 8 August 1996 claims 1-2	1-32
X WO 9743424 A1 (GENENCOR INTERNA 20 November 1997 (20.11.97) 21-34, page 12, lines 15-19	TIONAL, INC.), , see page 3, lines	1-8
A		7-32
X WO 9741213 A1 (NOVO NORDISK A/S (06.11.97), see page 11, li	), 6 November 1997 nes 5-12	1-8
А		9-32
	,	
X Further documents are listed in the continuation of Bo	x C. X See patent famil	y annex.
Special categories of cited documents:     A document defining the general state of the art which is not considered to be of particular relevance.	"T" later document published aft date and not in conflict with the principle or theory under	er the international filing date or prion:; the application but cited to understand iying the invention
"E" ether document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other	"X" document of particular relev	ance: the claimed invention cannot be be considered to involve an inventive
special reason (as specified)  "O" document referring to an oral disclosure, use, exhibition or other means	considered to involve an invo	ance: the claimed invention cannot be enaive step when the document is other such documents, such combination illed in the art
"P" document published prior to the international filing date but later that the priority date claimed	being obvious to a person sk  "&" document member of the sar	
Date of the actual completion of the international search	Date of mailing of the interna	ational search report
8 June 1999	0 8 -06- 19	99
Name and mailing address of the ISA/	Authorized officer	
Swedish Patent Office   Box 5055, S-102 42 STOCKHOLM	Yvonne Siösteen	
Facsimile No. + 46 8 666 02 86	Telephone No. + 46 8 782	25 00

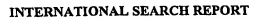
Facsimile No. +46 8 666 02 86
Form PCT/ISA/210 (second sheet) (July 1992)

## INTERNATIONAL SEARCH REPORT

International application No.

PCT/DK 99/00088

	1	
C (Continu	ation). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passage	ges Relevant to claim No.
A	Databas Swissprot, accessionno. P19531, Diderichsen B. et al: "Maltogenicalpha- amylase precursor (EC 3.2.1.133) Glucan 1,4-alpha-maltohydrolase)"; & Fems Microbiol. Lett. 56:53-60 (1988)	1-32
	<del></del>	
A	Dialog Information Services, file 5, Biosis Previews, Dialog accession no. 07281496, Biosis accession no. 000090061383, Boel E et al: "Calcium binding in alpha amylases an X-ray diffraction study at 2.1-Aresolution of two enzymes from aspergillus"; & Biochemistry 29 (26). 1990. 6244-6249	1-32
	~-	
A	Dialog Information Services, file 5, Biosis Previews, Dialog accession no. 09764348, Biosis accession no. 199598219266, Machius M et "Crystal structure of calcium-depleted Bacillus licheniformis alpha-amylase at 2.2 A resolution" Journal of Molecular Biology 246 (4):p545-559 19	; &
	<del></del>	
A	J. Mol. Biol., Volume 229, 1993, Changsoo Chang et al, "Crystallization and Preliminary X-ray Crystallographic Analysis of alpha-Amylase from Bacillus subtilis" page 235 - page 238	1-32
A	Plant Molecular Biology, Volume 25, 1994, Birte Svensson, "Protein engineering in the alpha-amylase family: catalytic mechanism, substrate specificity, and stability" page 141 - page 157	1-32



Information on patent family members

International application No.

03/05/99 | PCT/DK 99/00088

	atent document in search repor	rt	Publication date		Patent family member(s)		Publication date
WO	9623874	A1	08/08/96	AU BR CA CN EP JP	4483496 9607013 2211316 1172501 0808363 11500003	A A A	21/08/96 28/10/97 08/08/96 04/02/98 26/11/97 06/01/99
WO	9743424	A1	20/11/97	AU US	2996997 5763385		05/12/97 09/06/98
WO	9741213	A1	06/11/97	AU EP	2692897 0904360		19/11/97 31/03/99